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(54) Title: LIPOPEPTIDES AS ANTIBACTERIAL AGENTS

(57) Abstract: The present invention relates to novel lipopeptide compounds. The invention also relates to pharmaceutical compositions of these compounds and methods of using these compounds as antibacterial compounds. The invention also relates to methods of producing these novel lipopeptide compounds and intermediates used in producing these compounds.

LIPOPEPTIDES AS ANTIBACTERIAL AGENTS

FIELD OF THE INVENTION

The present invention relates to novel lipopeptide compounds. The invention also relates to pharmaceutical compositions of these compounds and methods of using these compounds as antibacterial compounds. The invention also relates to methods of producing these novel lipopeptide compounds and intermediates used in producing these compounds.

BACKGROUND OF THE INVENTION

The rapid increase in the incidence of gram-positive infections—including those caused by resistant bacteria—has sparked renewed interest in the development of novel classes of antibiotics. A class of compounds which have shown potential as useful antibiotics includes the A-21978C lipopeptides described in, for example, United States Patents RE 32,333; RE 32,455; RE 32,311; RE 32,310; 4,482,487; 4,537,717; and 5,912,226. Daptomycin, a member of this class, has potent bactericidal activity in vitro and in vivo against clinically relevant gram-positive bacteria that cause serious and life-threatening diseases. These bacteria include resistant pathogens, such as vancomycin-resistant enterococci (VRE), methicillin-resistant Staphylococcus aureus (MRSA), glycopeptide intermediate susceptible Staphylococcus aureus (GISA), coagulase-negative staphylococci (CNS), and penicillin-resistant Streptococcus pneumoniae (PRSP), for which there are few therapeutic alternatives. See, e.g., Tally et al., 1999, Exp. Opin. Invest. Drugs 8:1223-1238.

Despite the promise that antibacterial agents such as daptomycin offer, the need for novel antibiotics continues. Many pathogens have been repeatedly exposed to commonly-used antibiotics. This exposure has led to the selection of variant antibacterial strains resistant to a broad spectrum of antibiotics. The loss of potency and effectiveness of an antibiotic caused by resistant mechanisms renders the

antibiotic ineffective and consequently can lead to life-threatening infections that are virtually untreatable. As new antibiotics come to market pathogens may develop resistance or intermediate resistance to these new drugs, effectively creating a need for a stream of new antibacterial agents to combat these emerging strains. In addition compounds that exhibit bacteriacidal activity would offer advantages over present bacteriastatic compounds. Thus, novel synthetic antibacterial agents would be expected to be useful to treat not only "natural" pathogens, but also intermediate drug resistant and drug resistant pathogens because the pathogen has never been exposed to the novel antibacterial agent. Additionally, new antibacterial agents may exhibit differential effectiveness against different types of pathogens.

SUMMARY OF THE INVENTION

The present invention addresses this problem by providing novel lipopeptide compounds which have antibacterial activity against a broad spectrum of bacteria, including drug-resistant bacteria. Further, the compounds of the present invention exhibit bacteriacidal activity.

The present invention comprises, in one aspect, antibacterial compounds of Formula I:

and salts thereof,

wherein R is:

wherein X and X" are independently selected from C=O, C=S, C=NH, $C=NR^{X}$, S=O or SO₂;

wherein n is 0 or 1;

wherein R^X is selected from alkyl, alkenyl, alkynyl, aryl, heteroaryl, cycloalkyl, heterocyclyl, hydroxyl, alkoxy, carboxy or carboalkoxy;

wherein B is $X^{"}R^{Y}$, H, alkyl, alkenyl, alkynyl, aryl, heteroaryl, cycloalkyl or heterocyclyl,

wherein RY is selected from hydrido, alkyl, alkenyl, alkynyl, aryl, heteroaryl, cycloalkyl, heterocyclyl or hydroxyl,

wherein A is H, NH₂, NHR^A, NR^AR^B, alkyl, alkenyl, alkynyl, alkoxy, aryloxy, aryl, heteroaryl, cycloalkyl or heterocyclyl;

wherein R^A and R^B are independently selected from alkyl, alkenyl, alkynyl, aryl, heteroaryl, cycloalkyl, heterocyclyl or carboalkoxy;

wherein when n is 0, then A is additionally selected from:

$$- \begin{cases} -P & OR^{50} \\ OR^{51} & R^{52} \end{cases} \quad \text{and} \quad - \begin{cases} -P & OR^{50} \\ R^{53} & R^{53} \end{cases}$$

wherein each of R⁵⁰-R⁵³ is independently selected from C₁-C₁₅ alkyl, alternatively, wherein B and A together form a 5-7 membered heterocyclic or heteroaryl ring.

Wherein R¹ is

wherein X' and X"' are independently selected from C=O, C=S, C=NH, $C=NR^{X'}$, S=O or SO₂;

wherein m is 0 or 1;

wherein R^{X'} is selected from alkyl, alkenyl, alkynyl, aryl, heteroaryl, cycloalkyl, heterocyclyl, hydroxyl, alkoxy, carboxy or carboalkoxy;

wherein B' is $X'''R^{Y'}$, H, alkyl, alkenyl, alkynyl, aryl, heteroaryl, cycloalkyl or heterocyclyl; and

wherein R^Y is selected from hydrido, alkyl, alkenyl, alkynyl, aryl, heteroaryl, cycloalkyl, heterocyclyl or hydroxyl.

In one aspect of the invention, A' is H, NH₂, NHR^{A'}, NR^{A'}R^{B'}, heteroaryl, cycloalkyl or heterocyclyl,

wherein R^{A'} and R^{B'} are independently selected from alkyl, alkenyl, alkynyl, aryl, heteroaryl, cycloalkyl, heterocyclyl or carboalkoxy,

wherein when m is 0, then A' is additionally selected from:

wherein each of R^{50} - R^{53} is independently selected from C_1 - C_{15} alkyl; provided that when B' is H and X' is C=O, then A' is other than

- (a) a pyridinyl ring substituted with one substitutent NHC(O)R^D or
- (b) a C_5 - C_6 saturated cycloalkyl ring substituted with one substitutent NHC(O) R^D ;

wherein R^D is C_1 - C_{17} unsubstituted alkyl or C_2 - C_{17} unsubstituted alkenyl, and when B' is H and m=0, then A' is not H.

In another aspect of the invention, A' is aryl;

provided that when B' is H and X' is C=O, then A' is other than a phenyl ring substituted with substitutent NHC(O)R^D, wherein R^D is defined as above, which may be further optionally substituted on the phenyl ring with 1-2 substituents independently selected from amino, nitro, C_1 - C_3 alkyl, hydroxyl, C_1 - C_3 alkoxy, halo, mercapto, C_1 - C_3 alkylthio, carbamyl or C_1 - C_3 alkyl carbamyl.

In a third aspect of the invention, A' is alkyl, alkenyl, alkynyl, alkoxy or aryloxy;

provided that when B' is H and X' is C=O, then A' is other than

- (a) $-(C_1-C_{16} \text{ unsubstituted alkyl})-NH_2$;
- (b) -(C₁-C₁₀ unsubstituted alkyl)-NHC(O)R^D, wherein R^D is defined as described above:
- (c) $-C_1-C_{18}$ alkyl, optionally substituted with up to one hydroxyl, carboxyl or C_1-C_3 alkoxy, or one to three halo substituents;
 - (d) -C₄-C₁₈ unsubstituted alkenyl;

wherein R⁵⁴ is selected from C₁-C₁₇- unsubstituted alkyl or C₂-C₁₇- unsubstituted alkenyl; wherein R⁵⁵ is selected from hydroxyethyl, hydroxymethyl, mercaptomethyl, mercaptoethyl, methylthioethyl, 2-thienyl, 3-indolemethyl, phenyl optionally substituted with a group selected from halo, nitro, C₁-C₃-unsubstituted alkyl, hydroxy, C₁-C₃-unsubstituted alkoxy, C₁-C₃-unsubstituted alkylthio, carbamyl or C₁-C₃ unsubstituted alkylcarbamyl; or benzyl optionally substituted with a group selected from halo, nitro, C₁-C₃-unsubstituted alkyl, hydroxy, C₁-C₃-unsubstituted alkylthio, carbamyl or C₁-C₃ unsubstituted alkylcarbamyl; wherein t is 0 or 1 and wherein u is an integer from 1-3; and

when B' is H and X' is C=O, then X', together with A', does not form a carbamate amino protecting group, and

when B' is H and m is 0, then A' is other than C_4 - C_{14} unsubstituted alkyl.

In a fourth aspect of the invention, B' and A' together form a 5-7 membered heterocyclic or heteroaryl ring.

Wherein R² is

wherein K and K' together form a C_3 - C_7 cycloalkyl or heterocyclyl ring or a C_5 - C_{10} aryl or heteroaryl ring;

wherein J is selected from the group consisting of hydrido, amino, NHR^J, NR^JR^K, alkyl, alkenyl, alkynyl, alkoxy, aryloxy, aryl, heteroaryl, cycloalkyl, heterocyclyl, alkylamino, hydroxyl, thio, alkylthio, alkenylthio, sulfinyl, sulfonyl, azido, cyano, halo,

$$- \begin{cases} S \\ NR^{24}R^{25} \end{cases} \text{ and } - \begin{cases} S \\ OR^{26} \end{cases}$$

wherein each of R²⁴, R²⁵, and R²⁶ is independently selected from the group consisting of alkyl, cycloalkyl, heterocyclyl, aryl and heteroaryl; or R²⁴ and R²⁵ together form a 5-8 membered heterocyclyl ring;

wherein R^I and R^K are independently selected from alkyl, alkenyl, alkynyl, aryl, heteroaryl, cycloalkyl or heterocyclyl; or

alternatively, wherein J, together with R¹⁷, forms a 5-8 membered heterocyclyl or cycloalkyl ring; or

alternatively, wherein J, together with both R¹⁷ and R¹⁸, forms a 5-8 membered aryl, cycloalkyl, heterocyclyl or heteroaryl ring, and

wherein each of R¹⁷ and R¹⁸ is independently selected from the group consisting of hydrido, halo, hydroxyl, alkoxy, amino, thio, sulfinyl, sulfonyl and

wherein R^{17} and R^{18} taken together can form a group consisting of ketal, thioketal,

$$= \begin{cases} = 0 , = \\ = s , = \\ = NOR^{22} \text{ and } = \\ = NNR^{22}R^{23} \end{cases}$$

wherein each of R^{22} and R^{23} is independently selected from the group consisting of hydrido and alkyl.

In another embodiment, the invention also provides pharmaceutical compositions comprising compounds of Formula I and methods of use thereof.

In a further embodiment, the invention provides methods of making compounds of Formula I and pharmaceutical compositions thereof.

In a further embodiment, the invention provides compounds useful as intermediates for the preparation of compounds of Formula I.

In a still further embodiment, the invention provides methods of use of the compounds of Formula I to treat bacterial infections in humans.

DETAILED DESCRIPTION OF THE INVENTION

Definitions

Molecular terms, when used in this application, have their common meaning unless otherwise specified.

The term "hydrido" denotes a single hydrogen atom (H).

The term "acyl" is defined as a carbonyl radical attached to an alkyl, alkenyl, alkynyl, cycloalkyl, heterocycyl, aryl or heteroaryl group, examples including, without limitation, such radicals as acetyl and benzoyl.

The term "amino" denotes a nitrogen radical containing two substituents independently selected from the group consisting of hydrido, alkyl, cycloalkyl, carboalkoxy, heterocyclyl, aryl, heteroaryl and sulfonyl. Subsets of the

term amino are (1) the term "unsubstituted amino" which denotes an NH₂ radical, (2) the term "mono substituted amino" which is defined as a nitrogen radical containing a hydrido group and a substituent group selected from alkyl, cycloalkyl, heterocyclyl, aryl, or heteroaryl, and (3) the term "disubstituted amino" which is defined as a nitrogen radical containing two substituent groups independently selected from, alkyl, cycloalkyl, heterocyclyl, aryl, and heteroaryl. Preferred mono substituted amino radicals are "lower mono substituted amino" radicals, whereby the substituted amino radicals are "lower disubstituted amino" radicals, whereby the substituted amino" radicals, whereby the substituted

The term "acyloxy" denotes an oxygen radical adjacent to an acyl group.

The term "acylamino" denotes a nitrogen radical adjacent to an acyl group.

The term "carboalkoxy" is defined as a carbonyl radical adjacent to an alkoxy or aryloxy group.

The term "carboxyamido" denotes a carbonyl radical adjacent to an amino group.

The term "halo" is defined as a bromo, chloro, fluoro or iodo radical.

The term "thio" denotes a radical containing a substituent group independently selected from hydrido, alkyl, cycloalkyl, heterocyclyl, aryl and heteroaryl, attached to a divalent sulfur atom, such as, methylthio and phenylthio.

The term "alkyl" is defined as a linear or branched, saturated radical having one to about twenty carbon atoms unless otherwise specified. Preferred alkyl radicals are "lower alkyl" radicals having one to about five carbon atoms. One or more hydrogen atoms can also be replaced by a substitutent group selected from acyl, amino, acylamino, acyloxy, carboalkoxy, carboxy, carboxyamido, cyano, halo, hydroxyl, nitro, thio, alkyl, alkenyl, alkynyl, cycloalkyl, heterocyclyl, aryl, heteroaryl, alkoxy, aryloxy, sulfinyl, sulfonyl, oxo, guanidino, formyl and an amino acid side chain. Examples of alkyl groups include, without limitation, methyl, *tert*-butyl, isopropyl, and methoxymethyl. Subsets of the term alkyl are (1) "unsubstituted alkyl" which is defined as an alkyl group that bears no substituent groups (2) "substituted

alkyl" which denotes an alkyl radical in which (a) one or more hydrogen atoms is replaced by a substitutent group selected from acyl, acyloxy, carboalkoxy, carboxy, carboxyamido, cyano, nitro, thio, alkyl, alkenyl, alkynyl, cycloalkyl, heterocyclyl, aryl, heteroaryl, alkoxy, aryloxy, sulfinyl, sulfonyl, N-sulfonylcarboxyamido, N-acylaminosulfonyl or (b) two or more hydrogen atoms are each replaced by a substituent group independently selected from hydroxyl, carboxy, C₁-C₃ alkoxy, amino, acylamino, oxo or guanidino; and (3) the term "selected substituted alkyl" which denotes an alkyl radical in which (a) one proton is replaced by a group selected from hydroxyl, carboxy C₁-C₃ alkoxy, unsubstituted amino, acylamino, or acylamino phenyl or (b) one to three protons is replaced by a halo substituent.

The term "alkenyl" is defined as linear or branched radicals having two to about twenty carbon atoms, preferably three to about ten carbon atoms, and containing at least one carbon-carbon double bond. One or more hydrogen atoms can also be replaced by a substituent group selected from acyl, amino, acylamino, acyloxy, carboxy, carboxy, carboxyamido, cyano, halo, hydroxyl, nitro, thio, alkyl, alkenyl, alkynyl, cycloalkyl, heterocyclyl, aryl, heteroaryl, alkoxy, aryloxy, sulfinyl, sulfonyl, formyl, oxo and guanidino. The double bond portion(s) of the unsaturated hydrocarbon chain may be either in the cis or trans configuration. Examples of alkenyl groups include, without limitation, ethylenyl or phenyl ethylenyl.

The term "alkynyl" denotes linear or branched radicals having from two to about ten carbon atoms, and containing at least one carbon-carbon triple bond. One or more hydrogen atoms can also be replaced by a substituent group selected from acyl, amino, acylamino, acyloxy, carboalkoxy, carboxy, carboxyamido, cyano, halo, hydroxyl, nitro, thio, alkyl, alkenyl, alkynyl, cycloalkyl, heterocyclyl, aryl, heteroaryl, alkoxy, aryloxy, sulfinyl, sulfonyl, formyl, oxo and guanidino. An example of alkynyl group includes, without limitation, propynyl.

The term "aryl" or "aryl ring" denotes aromatic radicals in a single or fused carbocyclic ring system, having from five to fourteen ring members. In a preferred embodiment, the ring system has from six to ten ring members. One or more hydrogen atoms may also be replaced by a substituent group selected from acyl, amino, acylamino, acyloxy, azido, alkylthio, carboalkoxy, carboxy, carboxyamido,

cyano, halo, hydroxyl, nitro, thio, alkyl, alkenyl, alkynyl, cycloalkyl, heterocyclyl, aryl, heteroaryl, alkoxy, aryloxy, sulfinyl, sulfonyl and formyl. Examples of aryl groups include, without limitation, phenyl, naphthyl, biphenyl, terphenyl. Subsets of the term aryl are (1) the term "phenyl" which denotes a compound of the formula.

(2) the term "substituted phenyl" which is defined as a phenyl radical in which one or more protons are replaced by a substituent group selected from acyl, amino, acyloxy, azido, alkylthio, carboalkoxy, carboxy, carboxyamido, cyano, halo, hydroxyl, nitro, thio, alkyl, alkenyl, alkynyl, cycloalkyl, heterocyclyl, aryl, heteroaryl, alkoxy, aryloxy, sulfinyl, sulfonyl, N-sulfonylcarboxyamido, and N-acylaminosulfonyl and (3) the term "acylamino phenyl" denotes a phenyl radical in which one hydrogen atom is replaced by an acylamino group. One or more additional hydrogen atoms can also be replaced by a substituent group selected from acyl, amino, acylamino, acyloxy, azido, alkylthio, carboalkoxy, carboxy, carboxyamido, cyano, halo, hydroxyl, nitro, thio, alkyl, alkenyl, alkynyl, cycloalkyl, heterocyclyl, aryl, heteroaryl, alkoxy, aryloxy, sulfinyl, sulfonyl, N-sulfonylcarboxyamido, and N-acylaminosulfonyl.

"Heteroaryl" or "heteroaryl ring" denotes an aromatic radical which contain one to four hetero atoms or hetero groups selected from O, N, S,

from five to fifteen ring members. In a preferred embodiment, the heteroaryl ring system has from six to ten ring members. One or more hydrogen atoms may also be replaced by a substituent group selected from acyl, amino, acylamino, acyloxy, carboalkoxy, carboxy, carboxyamido, cyano, halo, hydroxyl, nitro, thio, thiocarbonyl, alkyl, alkenyl, alkynyl, cycloalkyl, heterocyclyl, aryl, heteroaryl, alkoxy, aryloxy, sulfinyl, sulfonyl, and formyl. Examples of heteroaryl groups include, without limitation, pyridinyl, thiazolyl, thiadiazoyl, isoquinolinyl, pyrazolyl, oxazolyl,

oxadiazoyl, triazolyl, and pyrrolyl groups. Subsets of the term heteroaryl are (1) the term "pyridinyl" which denotes compounds of the formula:

(2) the term "substituted pyridinyl" which is defined as a pyridinyl radical in which one or more protons is replaced by a substituent group selected from acyl, amino, acyloxy, carboalkoxy, carboxy, carboxyamido, cyano, halo, hydroxyl, nitro, thio, alkyl, alkenyl, alkynyl, cycloalkyl, heterocyclyl, aryl, heteroaryl, alkoxy, aryloxy, sulfinyl, sulfonyl, N-sulfonylcarboxyamido, and N-acylaminosulfonyl and (3) the term "acylamino pyridinyl" which denotes a pyridinyl radical in which one hydrogen atom is replaced by an acylamino group, additionally, one or more additional hydrogen atoms can also be replaced by a substituent group selected from acyl, amino, acylamino, acyloxy, carboalkoxy, carboxy, carboxyamido, cyano, halo, hydroxyl, nitro, thio, thiocarbonyl, alkyl, alkenyl, alkynyl, cycloalkyl, heterocyclyl, aryl, heteroaryl, alkoxy, aryloxy, sulfinyl, sulfonyl, N-sulfonylcarboxyamido, and N-acylaminosulfonyl.

The term "cycloalkyl" or "cycloalkyl ring" is defined as a saturated or partially unsaturated carbocyclic ring in a single or fused carbocyclic ring system having from three to twelve ring members. In a preferred embodiment, a cycloalkyl is a ring system having three to seven ring members. One or more hydrogen atoms may also be replaced by a substituent group selected from acyl, amino, acylamino, acyloxy, carboalkoxy, carboxy, carboxyamido, cyano, halo, hydroxyl, nitro, thio, alkyl, alkenyl, alkynyl, cycloalkyl, heterocyclyl, aryl, heteroaryl, alkoxy, aryloxy, sulfinyl, sulfonyl and formyl. Examples of a cycloalkyl group include, without limitation, cyclopropyl, cyclobutyl, cyclohexyl, and cycloheptyl.

The term "heterocyclyl," "heterocyclic" or "heterocyclyl ring" is defined as a saturated or partially unsaturated ring containing one to four hetero atoms

or hetero groups selected from O, N, NH,
$$-\xi - \frac{R^z}{N} = \frac{1}{N}$$
, wherein R^z is as defined for

R^X, ö, S, or on in a single or fused heterocyclic ring system having from three to twelve ring members. In a preferred embodiment, a heterocyclyl is a ring system having three to seven ring members. One or more hydrogen atoms may also be replaced by a substituent group selected from acyl, amino, acylamino, acyloxy, oxo, thiocarbonyl, imino, carboalkoxy, carboxy, carboxyamido, cyano, halo, hydroxyl, nitro, thio, alkyl, alkenyl, alkynyl, cycloalkyl, heterocyclyl, aryl, heteroaryl, alkoxy, aryloxy, sulfinyl, sulfonyl and formyl. Examples of a heterocyclyl group include, without limitation, morpholinyl, piperidinyl, and pyrrolidinyl.

The term "alkoxy" denotes oxy-containing radicals substituted with an alkyl, cycloalkyl or heterocyclyl group. Examples include, without limitation, methoxy, tert-butoxy, benzyloxy and cyclohexyloxy.

The term "aryloxy" denotes oxy-containing radicals substituted with an aryl or heteroaryl group. Examples include, without limitation, phenoxy.

The term "amino acid side chain" denotes any side chain (R group) from a naturally-occurring or a non-naturally occurring amino acid.

The term "sulfinyl" is defined as a tetravalent sulfur radical substituted with an oxo substituent and a second substituent selected from the group consisting of alkyl, cycloalkyl, heterocyclyl, aryl, or heteroaryl group.

The term "sulfonyl" is defined as a hexavalent sulfur radical substituted with two oxo substituents and a third substituent selected from alkyl, cycloalkyl, heterocyclyl aryl, or heteroaryl.

The term "carbamate amino protecting group" is defined as a recognized amino protecting group that when bound to an amino group forms a carbamate. Examples of carbamate amino protecting groups can be found in "Protective Groups in Organic Synthesis" by Theodora W. Greene, John Wiley and Sons, New York, 1981. Examples of carbamate amino protecting groups include benzyloxycarbonyl, t-butoxycarbonyl, t-amyloxycarbonyl, isobornyloxycarbonyl, adamantyloxycarbonyl, chlorobenzyloxycarbonyl, nitrobenzyloxycarbonyl or the like.

The salts of the compounds of the invention (preferably a compound of Formula I) include acid addition salts and base addition salts. In a preferred embodiment, the salt is a pharmaceutically acceptable salt of the compound of Formula I. The term "pharmaceutically-acceptable salts" embraces salts commonly used to form alkali metal salts and to form addition salts of free acids or free bases. The nature of the salt is not critical, provided that it is pharmaceutically-acceptable. Suitable pharmaceutically-acceptable acid addition salts of the compounds of the invention (preferably a compound of Formula I) may be prepared from an inorganic acid or an organic acid. Examples of such inorganic acids include, without limitation, hydrochloric, hydrobromic, hydroiodic, nitric, carbonic, sulfuric and phosphoric acid. Appropriate organic acids may be selected from aliphatic, cycloaliphatic, aromatic, arylaliphatic, heterocyclic, carboxylic and sulfonic classes of organic acids, examples of which include, without limitation, formic, acetic, propionic, succinic, glycolic, gluconic, maleic, embonic (pamoic), methanesulfonic, ethanesulfonic, 2hydroxyethanesulfonic, pantothenic, benzenesulfonic, toluenesulfonic, sulfanilic, mesylic, cyclohexylaminosulfonic, stearic, algenic, \(\beta\text{-hydroxybutyric, malonic,} \) galactic, and galacturonic acid. Suitable pharmaceutically-acceptable base addition salts of compounds of the invention (preferably a compound of Formula I) include. but are not limited to, metallic salts made from aluminum, calcium, lithium, magnesium, potassium, sodium and zinc or organic salts made from N,N'dibenzylethylenediamine, chloroprocaine, choline, diethanolamine, ethylenediamine, N-methylglucamine, lysine and procaine. All of these salts may be prepared by conventional means from the corresponding compound of the invention (preferably a compound of Formula I) by treating, for example, the compound of the invention (preferably a compound of Formula I) with the appropriate acid or base.

The compounds of the invention (preferably compounds of Formula I) can possess one or more asymmetric carbon atoms and are thus capable of existing in the form of optical isomers as well as in the form of racemic or non-racemic mixtures thereof. The compounds of the invention (preferably compounds of Formula I) can be utilized in the present invention as a single isomer or as a mixture of stereochemical isomeric forms. Diastereoisomers, i.e., nonsuperimposable stereochemical isomers,

can be separated by conventional means such as chromatography, distillation, crystallization or sublimation. The optical isomers can be obtained by resolution of the racemic mixtures according to conventional processes, for example by formation of diastereoisomeric salts by treatment with an optically active acid or base. Examples of appropriate acids include, without limitation, tartaric, diacetyltartaric, dibenzoyltartaric, ditoluoyltartaric and camphorsulfonic acid. The mixture of diastereomers can be separated by crystallization followed by liberation of the optically active bases from these salts. An alternative process for separation of optical isomers includes the use of a chiral chromatography column optimally chosen to maximize the separation of the enantiomers. Still another available method involves synthesis of covalent diastereoisomeric molecules by reacting compounds of the invention (preferably compounds of Formula I) with an optically pure acid in an activated form or an optically pure isocyanate. The synthesized diastereoisomers can be separated by conventional means such as chromatography, distillation, crystallization or sublimation, and then hydrolyzed to obtain the enantiomerically pure compound. The optically active compounds of the invention (preferably compounds of Formula I) can likewise be obtained by utilizing optically active starting materials. These isomers may be in the form of a free acid, a free base, an ester or a salt.

The invention also embraces isolated compounds. An isolated compound refers to a compound which represents at least 10%, preferably at least 20%, more preferably at least 50% and most preferably at least 80% of the compound present in the mixture. In a preferred embodiment, the compound, a pharmaceutically acceptable salt thereof, or a pharmaceutical composition comprising the compound exhibits a detectable (i.e. statistically significant) antimicrobial activity when tested in conventional biological assays such as those described herein.

Lipopeptide Compounds

A compound of the formula (I):

and salts thereof,

wherein R is:

wherein X and X" are independently selected from C=O, C=S, C=NH, C=NR X , S=O or SO₂,

wherein n is 0 or 1;

wherein R^X is selected from alkyl, alkenyl, alkynyl, aryl, heteroaryl, cycloalkyl, heterocyclyl, hydroxyl, alkoxy, carboxy or carboalkoxy,

wherein B is $X''R^Y$, H, alkyl, alkenyl, alkynyl, aryl, heteroaryl, cycloalkyl or heterocyclyl;

wherein R^Y is selected from hydrido, alkyl, alkenyl, alkynyl, aryl, heteroaryl, cycloalkyl, heterocyclyl or hydroxyl,

wherein A is H, NH₂, NHR^A, NR^AR^B, alkyl, alkenyl, alkynyl, alkoxy, aryloxy, aryl, heteroaryl, cycloalkyl or heterocyclyl;

wherein R^A and R^B are independently selected from alkyl, alkenyl, alkynyl, aryl, heteroaryl, cycloalkyl, heterocyclyl or carboalkoxy;

wherein when n is 0, then A is additionally selected from:

wherein each of R⁵⁰-R⁵³ is independently selected from C₁-C₁₅ alkyl, alternatively, wherein B and A together form a 5-7 membered heterocyclic or heteroaryl ring.

Wherein R¹ is

wherein X' and X"' are independently selected from C=O, C=S, C=NH, C=NR $^{X'}$, S=O or SO₂,

wherein m is 0 or 1;

wherein R^X is selected from alkyl, alkenyl, alkynyl, aryl, heteroaryl, cycloalkyl, heterocyclyl, hydroxyl, alkoxy, carboxy or carboalkoxy;

wherein B' is X"'RY, H, alkyl, alkenyl, alkynyl, aryl, heteroaryl, cycloalkyl or heterocyclyl, and

wherein RY is selected from hydrido, alkyl, alkenyl, alkynyl, aryl, heteroaryl, cycloalkyl, heterocyclyl or hydroxyl.

In one aspect of the invention, A' is H, NH₂, NHR^{A'}, NR^{A'}R^{B'}, heteroaryl, cycloalkyl or heterocyclyl,

wherein $R^{A'}$ and $R^{B'}$ are independently selected from alkyl, alkenyl, alkynyl, aryl, heteroaryl, cycloalkyl, heterocyclyl or carboalkoxy;

wherein when m is 0, then A' is additionally selected from:

$$- \left\{ \begin{array}{c} O \\ - P \\ O \\ O \\ O \\ \end{array} \right\} O = \left\{ \begin{array}{c} O \\ - P \\ - P \\ - P \\ \end{array} \right\} O = \left\{ \begin{array}{c} O \\ - P \\ - P \\ - O \\ \end{array} \right\} O = \left\{ \begin{array}{c} O \\ - P \\ - P \\ - O \\ \end{array} \right\} O = \left\{ \begin{array}{c} O \\ - P \\ - P \\ - O \\ \end{array} \right\} O = \left\{ \begin{array}{c} O \\ - P \\ - P \\ - O \\ \end{array} \right\} O = \left\{ \begin{array}{c} O \\ - P \\ - P \\ - O \\ \end{array} \right\} O = \left\{ \begin{array}{c} O \\ - P \\ - P \\ - O \\ \end{array} \right\} O = \left\{ \begin{array}{c} O \\ - P \\ - P \\ - O \\ \end{array} \right\} O = \left\{ \begin{array}{c} O \\ - P \\$$

wherein each of R^{50} - R^{53} is independently selected from C_1 - C_{15} alkyl; provided that when B' is H and X' is C=O, then A' is other than

- (a) a pyridinyl ring substituted with one substitutent NHC(O)R^D or
- (b) a C_5 - C_6 saturated cycloalkyl ring substituted with one substitutent NHC(O) R^D ;

wherein R^D is C_1 - C_{17} unsubstituted alkyl or C_2 - C_{17} unsubstituted alkenyl; and when B' is H and m=0, then A' is not H.

In another aspect of the invention, A' is aryl;

provided that when B' is H and X' is C=O, then A' is other than a phenyl ring substituted with substitutent NHC(O)R^D, wherein R^D is defined as above, which may be further optionally substituted on the phenyl ring with 1-2 substituents independently selected from amino, nitro, C_1 - C_3 alkyl, hydroxyl, C_1 - C_3 alkoxy, halo, mercapto, C_1 - C_3 alkylthio, carbamyl or C_1 - C_3 alkyl carbamyl.

In a third aspect of the invention, A' is alkyl, alkenyl, alkynyl, alkoxy or aryloxy;

provided that when B' is H and X' is C=O, then A' is other than

- (a) $-(C_1-C_{16} \text{ unsubstituted alkyl})-NH_2$;
- (b) $-(C_1-C_{10} \text{ unsubstituted alkyl})-NHC(O)R^D$, wherein R^D is defined as described above;
- (c) $-C_1-C_{18}$ alkyl, optionally substituted with up to one hydroxyl, carboxyl or C_1-C_3 alkoxy, or one to three halo substituents;
 - (d) -C₄-C₁₈ unsubstituted alkenyl;

wherein R⁵⁴ is selected from C₁-C₁₇- unsubstituted alkyl or C₂-C₁₇unsubstituted alkenyl; wherein R⁵⁵ is selected from hydroxyethyl, hydroxymethyl,
mercaptomethyl, mercaptoethyl, methylthioethyl, 2-thienyl, 3-indolemethyl, phenyl
optionally substituted with a group selected from halo, nitro, C₁-C₃-unsubstituted
alkyl, hydroxy, C₁-C₃-unsubstituted alkoxy, C₁-C₃-unsubstituted alkylthio, carbamyl
or C₁-C₃ unsubstituted alkylcarbamyl; or benzyl optionally substituted with a group
selected from halo, nitro, C₁-C₃-unsubstituted alkyl, hydroxy, C₁-C₃-unsubstituted
alkoxy, C₁-C₃-unsubstituted alkylthio, carbamyl or C₁-C₃ unsubstituted alkylcarbamyl;
wherein t is 0 or 1 and wherein u is an integer from 1-3; and

when B is H and X is C=O, then X, together with A, does not form a carbamate amino protecting group; and

when B' is H and m is 0, then A' is other than C_4 - C_{14} unsubstituted alkyl.

In a fourth aspect of the invention, B' and A' together form a 5-7 membered heterocyclic or heteroaryl ring.

Wherein R² is

wherein K and K' together form a C₃-C₇ cycloalkyl or heterocyclyl ring or a C₅-C₁₀ aryl or heteroaryl ring;

wherein J is selected from the group consisting of hydrido, amino, NHR^J, NR^JR^K, alkyl, alkenyl, alkynyl, alkoxy, aryloxy, aryl, heteroaryl, cycloalkyl, heterocyclyl, alkylamino, hydroxyl, thio, alkylthio, alkenylthio, sulfinyl, sulfonyl, azido, cyano, halo,

$$- \begin{cases} S & \text{and} \\ - S & \text{OR}^{26} \end{cases}$$

wherein each of R²⁴, R²⁵, and R²⁶ is independently selected from the group consisting of alkyl, cycloalkyl, heterocyclyl, aryl and heteroaryl, or R²⁴ and R²⁵ together form a 5-8 membered heterocyclyl ring;

wherein R^J and R^K are independently selected from alkyl, alkenyl, alkynyl, aryl, heteroaryl, cycloalkyl or heterocyclyl; or

alternatively, wherein J, together with R¹⁷, forms a 5-8 membered heterocyclyl or cycloalkyl ring; or

alternatively, wherein J, together with both R^{17} and R^{18} , forms a 5-8 membered aryl, cycloalkyl, heterocyclyl or heteroaryl ring, and

wherein each of R¹⁷ and R¹⁸ is independently selected from the group consisting of hydrido, halo, hydroxyl, alkoxy, amino, thio, sulfinyl, sulfonyl and

wherein R¹⁷ and R¹⁸ taken together can form a group consisting of ketal, thioketal,

wherein each of R²² and R²³ is independently selected from the group consisting of hydrido and alkyl.

In a preferred embodiment of the invention, R is selected from

wherein each of R³, R⁴ R⁵, and R⁶ is independently selected from the group consisting of hydrido, alkyl, aryl, heterocyclyl and heteroaryl, and wherein R⁴⁴ is selected from the group consisting of alkyl, aryl, heterocyclyl and heteroaryl.

In a more preferred embodiment of the invention R is selected from

wherein R^{41} is selected from the group consisting of alkyl, aryl-substituted alkyl, substituted phenyl, heterocyclyl, optionally substituted (C_8 - C_{14})-straight

In an even more preferred embodiment of the invention, R is

$$R^3$$
 (C_8 - C_{13})-straight-chain alkyl R^3 (C_8 - C_{13})-straight-chain alkyl R^3 (C_8 - C_{13})-straight-chain alkyl R^3 , and R^4 , and R^5 , R^4 , R^5 , R^6 , R^6

wherein X^3 is chloro or trifluoromethyl and wherein q is 0 or 1.

In a preferred embodiment of the invention, R¹ is selected from the group consisting of:

wherein R^8 is selected from an amino acid side chain, wherein said amino acid side chain may be one that is naturally occurring or one that is not naturally occurring, wherein each of R^9 , R^{10} and R^{11} is selected from hydrido, alkyl, aryl, heterocyclyl and heteroaryl; wherein R^{12} is selected from the group consisiting of heterocyclyl, heteroaryl, aryl, and alkyl and wherein R^{13} is selected from (C_1-C_3) -alkyl and aryl.

In a more preferred embodiment of the invention, R¹ is selected from the group consisting of

wherein R⁸ is selected from tryptophan side chain and lysine side chain; wherein each of R¹⁰ and R¹¹ is independently selected from hydrido and alkyl; wherein R¹² is selected from imidazolyl, N-methylimidazolyl, indolyl, quinolinyl, benzyloxybenzyl,

and benzylpiperidenylbenzyl; and wherein X^4 is selected from fluoro and trifluoromethyl

In a preferred embodiment of R², J is selected from the group

consisting of hydrido, amino, azido and

; wherein R¹⁷ and R¹⁸

taken together form a group selected from the group consisting of ketal,

$$= \begin{cases} = 0 & \text{and} & = \end{cases} = NOR^{22}$$

alternatively, R^{17} is hydroxyl when R^{18} is hydrido. Alternatively, wherein J, together with R^{17} , forms a heterocyclyl ring.

In a more preferred embodiment of the invention, R² is selected from

wherein R¹⁷ and R¹⁸ taken together form a group selected from

$$= \begin{cases} = 0 & \text{and} & = \begin{cases} = NOR^{22} \\ & \text{wherein } R^{22} \text{ is selected from the group} \end{cases}$$

consisting of H and alkyl; wherein R¹⁹ is selected from the group consisting of

In an even more preferred embodiment of the invention R² is

Another aspect of the present invention provides compounds of formula (I), wherein R is selected from NHCO-[(C_6 - C_{14})-alkyl]CH₃, and R¹ and R² are selected from Table A below. More preferably, R is selected from NHCO-[(CH_2)₆₋₁₄]- CH_3 .

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HN NHBOC	O NH ₂
HN NH ₂ OH	O NH2
HN N Cbz	NH,
HN NH	O NH ₂
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HN	O NH2
HN NH ₂	O NHz
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<u>Table I</u> Table I provides exemplary compounds of Formula I:

Table I provides exemplary compounds of Formula I:					
Cpd#	R	R ¹	R ²	Mass Spec	Synth Ex #
1	NHCO(CH ₂) ₈ CH ₃	-}-N NCO21Bu NHCO21Bu	O NH ₂	1863	6
2	NHCO(CH ₂) ₈ CH ₃	-N NH	O NH,	1663	6
3	NHCO(CH ₂) ₈ CH ₃	NHSO₂Ph	NH2	1762	5
4	NHCO(CH ₂) ₈ CH ₃	HN N N	O NH ₂	1792	4
5	NHCO(CH ₂) ₈ CH ₃	HN H	PE PE	1694	4
6	NHCO(CH ₂) ₈ CH ₃	7.	D Z-	1722	4
7	NHCO(CH ₂) ₈ CH ₃	HN N	O NH ₂	1764	4
8	NHCO(CH₂) ₈ CH₃	₹ - 2 - 2 -	Î-	1720	4
9	NHCO(CH ₂) ₈ CH ₃	O N-NH HN H CO ₂ H	Ž-()-	1775	4
10	NHCO(CH ₂) ₈ CH ₃	₹ 0= ₹ }-	Ž	1740	2
. 11	NHCO(CH ₂) ₈ CH ₃	2		1775	2
12	NHCO(CH ₂) ₈ CH ₃	O NH2		1820	2
13	NHCO(CH ₂) ₈ CH ₃	O NH ₂ CH ₃	0	1755.	2
14	NHCO(CH ₂) ₈ CH ₃	HN CH3	O NH2	1755	2
15	NHCO(CH ₂) ₈ CH ₃	O NH ₂	O NH2	1771	2

		,			
16	NHCO(CH₂)8CH₃	HN OCH3	NH2	1771	2
17	NHCO(CH ₂) ₈ CH ₃	O NH ₂	O NH2	1775	2
18	NHCO(CH ₂) ₈ CH ₃	O NO ₂	O NH2	1812	36
19	NHCO(CH ₂) ₈ CH ₃	O NH₂ HN CO₂H	NH2	1785	2
20	NHCO(CH ₂) ₈ CH ₃	O NHCH,	N. T.	1755	2
21	NHCO(CH ₂) ₈ CH ₃	O OCH3	O NH2	1756	3Ъ
22	NHCO(CH ₂) ₈ CH ₃	P - OF	NH ₂	1757	2
23	NHCO(CH ₂) ₈ CH ₃	HN. NH.	O NH	1742	2
24	NHCO(CH ₂) ₈ CH ₃	O NH.	O NH ₂	1790	2
25	NHCO(CH ₂) ₈ CH ₃	₩ ₩ ₩ ₩ ₩ ₩ ₩ ₩ ₩ ₩ ₩ ₩ ₩ ₩ ₩ ₩ ₩ ₩ ₩	O NH ₂	1758	2
26	NHCO(CH ₂) ₈ CH ₃	NH2 HN2	0 NH2	1758	2
27	NHCO(CH ₂) ₈ CH ₃		0= \-\-\-\-\-\-\-\-\-\-\-\-\-\-\-\-\-\-\-	1758	2 .
28	NHCO(CH ₂) ₈ CH ₃	± }-	£	1726	3b
29	NHCO(CH ₂) ₈ CH ₃		0=\\-\-\-\-\-\-\-\-\-\-\-\-\-\-\-\-\-\-\	1728	3b
30	NHCO(CH ₂)gCH ₃	HN NH ₂		1741	3b
31	NHCO(CH ₂) ₈ CH ₃	HN NH2	£	1741	3b

32	NHCO(CH ₂) ₈ CH ₃	HN TO SUNS	O NH ₂	1771	3ъ
33	NHCO(CH ₂) ₈ CH ₃	HN-	O NH2	1851	3Ъ
34	NHCO(CH ₂) ₈ CH ₃	O N(CH ₃) ₂	O NH2	1767	3Ъ
35	NHCO(CH ₂) ₈ CH ₃	HN	NH ₂	1782	3Ъ
36	NHCO(CH ₂) ₈ CH ₃	O NHCH3	27	1780	8
37	NHĊO(CH₂)₅CH₃	HN S		1873	8
38	NHCO(CH ₂) ₈ CH ₃	HN F	O NH2	1729	1
39	NHCO(CH ₂) ₈ CH ₃	о ососн ₃ ни со₂н ососн ₃	O NH ₂	1838	3b
40	NHCO(CH ₂) ₈ CH ₃	och .	O NH ₂	1741	. 1
41	NHCO(CH₂)8CH₃	HN NHBOC	O NH ₂	1908	3
42	NHCO(CH₂)8CH₃	O HN CO₂CH₃ NHBOC	O NH2	1865	3
43	NHCO(CH ₂) ₈ CH ₃	HN CO2'Bu	NH ₂	1893	3
44	NHCO(CH ₂) ₈ CH ₃	HN NHBOC N	O NH ₂	1908	3
45	NHCO(CH ₂) ₈ CH ₃	HN-NH2 NH2	O NH2	1808	3
46	NHCO(CH ₂) ₈ CH ₃	HN CO ₂ CH ₃	O NH2	1764	3
47	NHCO(CH ₂) ₈ CH ₃	O HN CONH ₂	O NH2	1750	3
48	NHCO(CH ₂) ₈ CH ₃	HN CONH ₂	O NH ₂	1736	3

49	NHCO(CH ₂) ₈ CH ₃	HN NHBOC	O NH2	2004	3a
50	NHCO(CH ₂) ₈ CH ₃	HN	D =	1712	1
51	NHCO(CH ₂) ₈ CH ₃	HN NH ₂ NHTs	\$ - \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \	1904	3a
52	NHCO(CH ₂) ₈ CH ₃	HŅ CH₃	0=\\-\-\-\-\-\-\-\-\-\-\-\-\-\-\-\-\-\-\	1725	1
54	NHCO(CH ₂) ₈ CH ₃	HN NH ₂	D=	1749	3a
55	NHCO(CH ₂) ₈ CH ₃	HN NHBOC	E -	1884	3
56	NHCO(CH ₂) ₈ CH ₃	HN NH ₂ OH	0 ZH2	1785	3
57	NHCO(CH ₂) ₈ CH ₃	HN N Cbz	E	1853	3
58	NHCO(CH ₂) ₈ CH ₃	HN HN S	O NH ₂	1847	3
60	NHCO(CH ₂) ₈ CH ₃	HN NH2 NH2	O NH2	1778	3
61	NHCO(CH ₂) ₈ CH ₃	HN THE	O NH2	1792	3
- 62	NHCO(CH ₂) ₈ CH ₃	HN NH, NH	O NH ₂	1826	3
63	NHCO(CH ₂) ₈ CH ₃	HN NH ₂ NH ₃	O NH ₂	1826	3
64	NHCO(CH ₂) ₈ CH ₃	NH NH NH	O NH ₂	1838	. 3
65	NHCO(CH ₂) ₈ CH ₃	HN NH ₂ O NH ₂	D NH2	1812	3
66	NHCO(CH ₂) ₈ CH ₃	HN NH2 NH	0 = 1	1808	3

67	NHCO(CH ₂)8ĊH ₃	HN NH ₂	O NH2	1769	3
68	NHCO(CH ₂) ₈ CH ₃	HN NH ₂ S	O NH ₂	1824	3
69	NHCO(CH₂)₅CH₃	HN NH ₂ N N	NH2	1775	3
7 0	NHCO(CH ₂) ₈ CH ₃	HN HN N	O NH ₂	1820	3
72	NHCO(CH ₂) ₈ CH ₃	HN NH ₂	O NH?	1707	3
73.	NHCO(CH ₂) ₈ CH ₃	HN NH ₂ NN	O NH ₂	1758	3
74	NHCO(CH₂)8CH₃	HN NH ₂ N NBOC	O NH2	1959	-3
75	NHCO(CH ₂) ₈ CH ₃	. HN NHBOC	O NH ₂	1810	3
76	NHCO(CH ₂) ₈ CH ₃	CH2-	NH ₂	1787	1g
77	NHCO(CH ₂) ₈ CH ₃	NH(CH₂)₂OH	O NH ₂	1665	1
78	NHCO(CH ₂) ₈ CH ₃	HŅ NHPh N NH2	O NH ₂	1820	1
79	NHCO(CH ₂) ₈ CH ₃	NH NH	O NH ₂	1750	1
80	NHCO(CH ₂) ₈ CH ₃	NH NH OCH,	O No.	1779	1
81	NHCO(CH ₂) ₈ CH ₃	NH NH F	O NH ₂	1767	le
82	NHCO(CH ₂) ₈ CH ₃	NH NH CH ₃	O NH ₂	1763	1
83	NHCO(CH ₂) ₈ CH ₃	HN C	O NH2	1869	1
84	NHCO(CH₂)8CH₃	CH ₃	O NH ₂	1764	1

85	NHCO(CH ₂) ₈ CH ₃	HN CH3	O NH ₂	1714	1c
86	. HN NH2 NTS	HN NH ₂	O NH2	1935	9
87	NHCO(CH ₂) ₈ CH ₃	HN NO	O NH ₂	1863	1
88	NHCO(CH ₂) ₈ CH ₃	N CI Z	O NH ₂	2151	1
89	NHCO(CH ₂) ₈ CH ₃	HN CI	0 NH2	1887	1
90	NHCO(CH ₂) ₈ CH ₃	N O O OME	O NH ₂	2046	1
91	NHCO(CH ₂) ₈ CH ₃	N NE 12	O NH2	1996	1
92	NHCO(CH ₂) ₈ CH ₃	HN NEt2	O = ==================================	1809	. 1.
93	NHCO(CH ₂) ₈ CH ₃	HN O ⁿ Bu	O NH ₂	1783	1
94	NHCO(CH ₂) ₈ CH ₃	HN O ⁿ Pr	O NH ₂	1770	1
95	NHCO(CH ₂) ₈ CH ₃	HN C	O NH,	1836	1
96	NHCO(CH ₂) ₈ CH ₃	HN MeO	O NH2	1792	1
97	NHCO(CH ₂) ₈ CH ₃	OMe HN -	O NH2	1847	1
98	NHCO(CH ₂) ₈ CH ₃	N () 2	O NH ₂	1838	1
99	NHCO(CH ₂) ₈ CH ₃	N F	ž-	1837	1
100	NHCO(CH ₂) ₈ CH ₃	HN CO	0 NH ₂	1817	1
101	HN CI	HN NH2	O NH ₂	1867	9

100	NILICO/CH V CH	O NH	O NH ₂	1840	_
102	NHCO(CH ₂) ₁₁ CH ₃	HN NH ₂	+0	1849	9
103	NHCO(CH ₂) ₈ CH ₃		NH ₂	1885	i .
104	NHCO(CH ₂) ₈ CH ₃	× N N N 2	O NH2	2150	Î
105	NHCO(CH ₂) ₈ CH ₃	HN NO	O NH2	1756	1
106	NHCO(CH ₂) ₈ CH ₃	HN OH	O NH2	1833	1
107	NHCO(CH ₂) ₈ CH ₃	HN CF3	NH.	1871	1
108	NHCO(CH ₂) ₈ CH ₃	HN CI	O NH ₂	1873	1
109	NHCO(CH ₂) ₈ CH ₃	HN CI	O NH ₂	1872	1
110	NHCO(CH ₂) ₈ CH ₃	m (C) a C)	NH ₂	2014	1
111	NHCO(CH ₂) ₈ CH ₃	HW O O	O NH ₂	1817	1
112	NHCO(CH ₂) ₈ CH ₃	N CF5	O NH ₂	2121	1
. 113	NHCO(CH ₂) ₈ CH ₃	NE1	NH ₂	2036	1
114	NHCO(CH ₂) ₈ CH ₃	AN HEI	O NH ₂	1826	1
115	NHCO(CH ₂) ₈ CH ₃	HN NC	NH ₂	1736	I
116	NHCO(CH ₂) ₈ CH ₃	HN-	NH2 O	1797	1

117	NHCO(CH ₂) ₈ CH ₃	HN O Bu	E-	1860	1
118	NHCO(CH ₂) ₈ CH ₃	N CI)	ž O - -	2055	1
119	NHCO(CH ₂) ₈ CH ₃	HN^ CI	Î-	1837	1
120	NHCO(CH ₂) ₈ CH ₃	N (0 (NO ₂)	NHÎ -	2104	1
121	NHCO(CH ₂) ₈ CH ₃	HN °	O NH ₂	1803	1
122	NHCO(CH ₂) ₈ CH ₃	HN CO₂H	£	1755	1
123	NHCO(CH ₂) ₈ CH ₃	HN O'Hex	P. P.	1812	1
124	NHCO(CH ₂) ₈ CH ₃	N OnHex	NH2	2002	. 1
125	NHCO(CH ₂) ₈ CH ₃	N O'Bu	₹2 	1946	1
126	NHCO(CH ₂) ₈ CH ₃	N O ⁿ Pr	NH2	1918	1
127	NHCO(CH ₂) ₈ CH ₃	NH —	O NH ₂	1811	1
128	NHCO(CH ₂) ₈ CH ₃	N Coopf	O NH ₂	2050	· l
129	NHCO(CH ₂) ₈ CH ₃	HN NO ₂	O NH ₂	1756	1
130	NHCO(CH ₂) ₈ CH ₃	HIN CN	O NH ₂	1762	1
131	NHCO(CH ₂) ₈ CH ₃	N N N	O NH ₂	1904	1

132	NHCO(CH ₂) ₈ CH ₃	OMe 2		1962	1
133	NHCO(CH ₂) ₈ CH ₃	-\\\\-\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\	Ž O - -	1726	1
134	NHCO(CH ₂) ₈ CH ₃	OMe O O	D NH2	2074	1
135	NHCO(CH ₂) ₈ CH ₃	HN F	Ž-(-)	1729	1
136	NHCO(CH ₂) ₈ CH ₃	£	ž-(-)	1729	1
137	NHCO(CH ₂) ₈ CH ₃	N ()	Ž	2014	1
. 138	NHCO(CH ₂) ₈ CH ₃	TZ-	NH ₂	1762	1
139	NHCO(CH ₂) ₈ CH ₃	HN	O NH ₂	1751	1
140	NHCO(CH ₂) ₈ CH ₃	N CO	NH ₂	1881	1
141	NHCO(CH ₂) ₈ CH ₃	N ()	O NH ₂	1914	1
142	NHCO(CH ₂) ₈ CH ₃	HN	O NH	1753	1
143	NHCO(CH ₂) ₈ CH ₃	HN O	O NH ₂	1803	1
144	NHCO(CH ₂) ₈ CH ₃	HN	O NH ₂	1813	1
145	NHCO(CH ₂) ₈ CH ₃	N Phy 2	NH ₂	2006	1
146	NHCO(CH ₂) ₈ CH ₃	HN N	O NH2	1701	1
147	NHCO(CH ₂) ₈ CH ₃	HN	NH2	1799	1

148	NHCO(CH ₂) ₈ CH ₃		O NH ₂	19 78	1
149	NHCO(CH ₂) ₈ CH ₃	HATTON	D = \{-	1834	1
150	NHCO(CH ₂) ₈ CH ₃		O NET	1777	1
151	NHCO(CH ₂) ₈ CH ₃	HN OMe	Ž	1847	1
152	NHCO(CH ₂) ₈ CH ₃	N OMe O	Ž,	2074	1
153	NHCO(CH ₂) ₈ CH ₃	HN O ⁿ Dodecyl	Ĭ-	1895	. 1
154	NHCO(CH ₂) ₈ CH ₃	HN O ⁿ Decyl	NHT NHT	1867	1
155	NHCO(CH ₂) ₈ CH ₃	HN O ⁿ Octyl	O NH2	1839	1
156	NHCO(CH ₂) ₈ CH ₃	HŅ CO2H	O NH ₂	1781	1
157	NHCO(CH ₂) ₈ CH ₃	HN NMe ₂	NH ₂	1780	-1
158	NHCO(CH ₂) ₈ CH ₃	HŅ	O NH,	1781	1
159	NHCO(CH ₂) ₈ CH ₃	HN N-Ph	O NH ₂	1805	1
160	NHCO(CH ₂) ₈ CH ₃	N-Ph	O NH ₂	1990	1
161	NHCO(CH ₂) ₈ CH ₃	HN CO2H	O NH ₂	1785	1
162	NHCO(CH ₂) ₈ CH ₃	N Br) 2	O NH ₂	2092	1
163	NHCO(CH ₂) ₈ CH ₃	NO ₂	NH ₂	1944	1

164	NHCO(CH ₂) ₈ CH ₃	HN OOO	O NH ₂	1817	1
165	NHCO(CH ₂) ₈ CH ₃	N/ Ph)	O NH ₂	2014	1
166	NHCO(CH ₂) ₈ CH ₃	HN F	O NH ₂	1747	1
167	NHCO(CH ₂) ₈ CH ₃	N CO	0 NH2	1853	1
168	NHCO(CH ₂) ₈ CH ₃	HN Z	O N-	1762	1
169	NHCO(CH ₂) ₈ CH ₃	<u>N</u> () ₂	D E	1829	1
17 1	NHCO(CH ₂) ₈ CH ₃	N (Butyi)		1914	1
172	NHCO(CH ₂) ₈ CH ₃	HN n _{Butyl}	P P	1767	1
173	NHCO(CH ₂) ₈ CH ₃	HN	D = \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \	1736	1
174	NHCO(CH ₂) ₈ CH ₃	HN S	Ĩ- -	1718	1
175	NHCO(CH ₂) ₈ CH ₃	HN Pentyl	₽	1808	1
176	NHCO(CH ₂) ₈ CH ₃		NH2	1781	1
177	NH₂		₹	1632	1
178	NHCO(CH ₂) ₈ CH ₃	HN NH ₂	0	1783	3
179	NHCO(CH ₂) ₈ CH ₃	HN NH2 NHBOC	DE P	1884	3
180	NHCO(CH ₂) ₈ CH ₃	HN NHFmoc	NH ₂	1905	. 3

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181	NHCONH(CH₂)₁₀CH₃	HN NH ₂ N	O NH ₂	1851	9
182	NHCO(CH ₂) ₈ CH ₃	HN	NH2	1801	3ъ
183	NHCO(CH ₂) ₈ CH ₃	N OH)	O NH2	1833	1
184	NHCO(CH ₂) ₈ CH ₃	HNOH	NET.	1727	1 .
185	NHCO(CH ₂) ₈ CH ₃	HN OH	NHT NHT	1743	1
186	NHCO(CH ₂) ₈ CH ₃	N OO	NH ₂	1890	. 1
187	NHCO(CH ₂) ₈ CH ₃	HN	NH ₂	1756	3
189	NHCO(CH ₂) ₈ CH ₃	}-N	O NH2	1717	3b
190	NHCO(CH ₂) ₈ CH ₃	O SO ₃ H	O NH ₂	1805	2
192	NHCO(CH ₂) ₈ CH ₃	HN NH ₂ OH	2-{-} 	1811	8
193	NHCO(CH ₂) ₈ CH ₃	B S S S S S S S S S S S S S S S S S S S	N. T.	1836	3
194	NHCO(CH ₂) ₈ CH ₃	HN OCF3	0 =	1795	1
195	NHCO(CH ₂) ₈ CH ₃	OCF ₃	NH2	1862	1
196	NHCO(CH ₂) ₈ CH ₃	HN CI	2 × × × × × × × × × × × × × × × × × × ×	1780	1
197	NHCO(CH ₂) ₈ CH ₃	HN CI	£	1746	1
198	NHCO(CH ₂) ₈ CH ₃	HN N(CH ₃) ₂	DÎ -	1754	1

199	NHCO(CH ₂) ₈ CH ₃	HŅ CI	O NH2	1780	1
200	NHCO(CH ₂) ₈ CH ₃	HN NH ₂ NH		1792	8a
201	NHCO(CH ₂) ₈ CH ₃	HN	NE ₂	1821	1
202	NHCO(CH ₂) ₈ CH ₃	HN NMe ₂	NH ₂		1
203	NHCO(CH ₂) ₈ CH ₃	HN NH ₂ N	O NH2	1793	1
204	NHCO(CH ₂) ₈ CH ₃	HN NHBo c NH	O NH ₂	1893	
205	NH(CH ₂) ₈ CH ₃	HN NH ₂ NH	O NH ₂	1779	9a
206	NHCO(CH ₂) ₈ CO ₂ Me	HN NH ₂	O NH ₂	1851	9
207	NHCO(CH₂) ₆ CO₂Me	HN NH ₂ NH	O NET CONTRACTOR	1823	9
208	NHCO(CH ₂) ₈ CH ₃	HN Pr	\$-\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\	1878	1
209	NHCO(CH ₂) ₈ CH ₃	HN O ₂		1880	1h
210	NHCO(CH ₂) ₈ CH ₃	HN CI	O NH ₂	1851	1
211	NHCO(CH ₂) ₈ CH ₃	HN OBn	O NHÃ	1924	1
212	NHCO(CH ₂) ₈ CH ₃	HN N	NH ₂	1701	1d
213	NHCO(CH ₂) ₆ NHBoc	HN NHBo c NH	O NH2	1980	9

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214	NHCO(CH ₂) ₇ NHBoc	HN NHBo c N H	O NH ₂	1994	9
215	NHCO(CH ₂)₁₀NHBoc	HN NHBo c N H	O NH2	2036	9
216	NHCO(CH ₂) ₁₁ NHBoc	HN NHBo c NH	O NH2	2050	9
217	NHCO(CH ₂) ₁₀ NH ₂	HN NH ₂	O NH ₂	1836	9
218	NHCO(CH ₂) ₁₁ NH ₂	HN NH2 N	O NH ₂	1850	9
219	NHCO(CH ₂) ₆ CH(CH ₃) ₂	HN NH ₂ N H	O NH ₂	1807	9
220	NHCONH(CH ₂) ₁₁ CH ₃	HN NH ₂ N	NH ₂	1865	9
221	NHCO(CH ₂) ₈ CH ₃	I NI	O NH2	1807	6
222	NHCO(CH ₂) ₈ CH ₃		0 = \(\frac{1}{\frac{1}{\chi_{-}}} \)	1935	1
223	NHCO(CH ₂) ₈ CH ₃	HIN N	£	1779	1
224	NHCO(CH ₂) ₈ CH ₃	HN NHBoc NHBoc	-}- (1936	1
225	NHCO(CH ₂) ₈ CH ₃	HN NH ₂	NH2	1735	1
226	NHCO(CH ₂) ₈ CH ₃		O NH2	1958	1
227	NHCO(CH ₂) ₈ CH ₃	HW N-Ph	O NH ₂	1899	1
228	NHCO(CH ₂) ₈ CH ₃	HN N	O NH ₂	1917	1

.229	NHCO(CH ₂) ₈ CH ₃		NH ₂	1914	1
230	NHCO(CH ₂) ₈ CH ₃	HN - CI	O NH ₂	1969	1
231	NHCO(CH ₂) ₈ CH ₃	HN N Ph	O NH2	1990	1
232	NHCO(CH ₂) ₈ CH ₃	HN N Ph	O RE2	1940	1
233	NHCO(CH ₂) ₈ CH ₃	HN ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~	O NH ₂	1902	1
234	NHCO(CH ₂) ₈ CH ₃	HN N N	O NH2	1901	1
- 235	NHCO(CH ₂) ₈ CH ₃	HN N-C-CI	O NH2	1934	1
236	NHCO(CH₂)₃CH₃		NH2	1984	1
237	NHCO(CH ₂) ₈ CH ₃		O NH2	1926	1
238	NHCO(CH ₂) ₈ CH ₃	HN N-C-F	o T	1944	1
239	NHCO(CH ₂) ₈ CH ₃	HN N-Bn	Î-	1940	1
240	NHCO(CH ₂) ₈ CH ₃		Ĭ- - - -	1995	1
241	NHCO(CH ₂) ₈ CH ₃	HN Ph		2016	1
242	NHCO(CH ₂) ₈ CH ₃		D Z	1928	1
243	NHCO(CH ₂) ₈ CH ₃		O NH2	1927	1
244	NHCO(CH ₂) ₈ CH ₃	HV C C	0=\\-\-\-\-\-\-\-\-\-\-\-\-\-\-\-\-\-\-\	1960	1
245	NHCO(CH ₂) ₈ CH ₃	HN. NH2	D = 1	1790	3
246	HN CI	HN NH ₂ N	NH ₂	1807	9

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247	HN CI	HN NH ₂ N	NH ₂	1841	9
248	HN OPh	HIN NH2 NH	NH2	1864	9
249	HN O ⁿ Buty	HIV NH ₂ NH	ž.	1843	9
250	HN CI	HIN NH ₂ NH	0 =	1882	9
251	HN CI	HN NH ₂	- Z	1823	9
252	NHCO(CH ₂) ₈ CH ₃	HN NO ₂ N-Bn	0 -	1931	1
253	NHCO(CH ₂) ₈ CH ₃	HN N-Bn	O NH?	1886	1f
254	NHCO(CH ₂) ₇ CH ₃	NBoc HN NHBoc	O NH2	1650	7
255	NHCO(CH₂)₀CH₃	NBoc HN NHBoc	O NH2	1678	.7
256	NHCO(CH ₂) ₁₀ CH ₃	NBoc HN NHBoc	D N N N N N N N N N N N N N N N N N N N	1692	7
257	NHCO(CH ₂) ₁₁ CH ₃	NB& HIV NHBoc	Ĭ,	1706	7
-258	NHCO(CH ₂) ₁₂ CH ₃	NBoc HN NHBoc	D Z	1720	7a _.
259	NHCO(CH ₂) ₈ CH ₃	NBC NBC	≥	1706	6
260	NHCO(CH ₂) ₉ CH ₃	NH HN NH ₂	- -	1678	7
261	NHCO(CH ₂) ₁₁ CH ₃	NH HN NH ₂	0	1705	7

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262	NHCO(CH ₂) ₁₂ CH ₃	HN NH ₂	NH2	1719	7a
263	HN CI	NBoc HN NHBoc	O NH ₂	1738	7
264	N=N N-heptyl	HN NH, NH	O NH2	1862	9
265	NHCO(CH ₂) ₈ CH ₃	HN CI	O NH2	1890	1
266	NHCO(CH ₂) ₈ CH ₃	HN NO ₂	O NH2	1841	1
267	NHCO(CH ₂) ₈ CH ₃	HN S	O NH ₂	1910	· 1
268	NHCO(CH ₂) ₈ CH ₃	HIN N-N-CF3	O NH ₂	1940	9
269	N=N N-nHeptyl	NHB& ZH	O NH ₂	1862	6
270	NHCO(CH ₂) ₈ CH ₃	HV NH	O NH2	1706	6
271	NHCO(CH ₂) ₈ CH ₃	HH.	D = 1	1851	1
272	NHCO(CH ₂) ₈ CH ₃	N CI	DE NOTE OF THE PROPERTY OF THE	2081	1 .
273	NHCO(CH₂) ₈ CH ₃	N OMe		1964	1
274	NHCO(CH₂)₃CH₃	HN OMe	DE 2	1793	1
275	NHCO(CH ₂) ₈ CH ₃	HN CI	NET ?	1797	1
276	NHCO(CH ₂) ₈ CH ₃	N CI)	O NH2	1973	1

277	NHCO(CH ₂) ₈ CH ₃	HN C	O NH ₂	1778	1
278	NHCO(CH ₂) ₈ CH ₃	HIT L	O NH2	1780	1
279	NHCO(CH ₂) ₈ CH ₃	"(") ₂	O NH2	1940	1
280	NHCO(CH ₂) ₈ CH ₃	HIM CI	NH ₂	1797	1
281	NHCO(CH ₂) ₈ CH ₃	m Cl2	O NH2	1974	1
282	NHCO(CH ₂) ₈ CH ₃	HN NO ₂	O NH2	1807	la
283	NHCO(CH ₂) ₈ CH ₃	HN CI	NH ₂	1797	1
284	NHCO(CH ₂) ₈ CH ₃	N (N ())2	Ž- (-)	1973	1
285	NHCO(CH ₂) ₈ CH ₃	HP. C		1796	1b
286	NHCO(CH ₂) ₈ CH ₃	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	±	1898	1
287	NHCO(CH ₂) ₈ CH ₃	OWH,		1806	1
288	NHCO(CH ₂) ₈ CH ₃	HK CI	₹ 0 - -	1812	1
289	NHCO(CH ₂) ₈ CH ₃	HN N(CH ₂) ₂		1806	1
290	NHCO(CH ₂) ₈ CH ₃	NO ₂	O NE	1806	1

291	NHCO(CH₂)8CH₃	HIN NOH CI	O NH ₂	1848	1
292	HN CI	HN NH ₂	O NH2	1738	7
293	NHCO(CH ₂) ₁₀ CH ₃	NH HN NH ₂	O NH2	1692	7
294	NHCO(CH₂)₁CH₃	NH HN NH ₂	O NH ₂	1650	7
295	NHCO(CH ₂) ₁₁ CH ₃	HN NHBoc	D H	1991	10ъ
296	NHCO(CH ₂) ₁₀ CH ₃	HN NH Boc	NH2	1978	10b
297	NHCO(CH ₂) ₉ CH ₃	HN NH Boc	O NET	1964	10Ъ
298	NHCONH(CH₂)7CH₃	HN NH Boc	O NH.	1950	10ъ
299	NHCONH(CH ₂) ₁₀ CH ₃	HN NH Boc	O NET	1992	10b
300	NHCONH(CH ₂) ₁₁ CH ₃	HN NH Boc		2006	1 0 b
301	NHCO(CH ₂) ₁₁ CH ₃	HN NH ₂	Ĭ-	1791	10b
302	NHCO(CH ₂) ₁₀ CH ₃	HN NH ₂	Î	1778	10b
303	NHCO(CH₂)₀CH₃	HN NH ₂		1764	10b
304	NHCONH(CH ₂) ₇ CH ₃	HN NH ₂	D = 1	1750	10ъ
305	NHCONH(CH ₂) ₁₀ CH ₃	HN NH ₂	PE ?	1792	10ъ
306	NHCONH(CH ₂) ₁₁ CH ₃	HN NH ₂	O NH2	1806	10ъ

	,				
307	NHCO(CH ₂) ₉ CH ₃	HN NHBac N	O NH ₂	1922	10b
308	NHCO(CH ₂) ₁₀ CH ₃	HN NHBœ N	O NH2	1936	10b
3,09	NHCO(CH ₂) ₁₀ CH ₃	HN NH2 NH	O NE	1836	10b
310	NHCO(CH ₂) ₉ CH ₃	HN NH ₂ NH	O NH2	1821	10b
311	NHCONH(CH ₂) ₇ CH ₃	HN NH ₂ NH	O NH2	1808	10b
312	NHCONH(CH₂)⁊CH₃	HN F NH2	O NH ₂	1759	10ь
313	NHCONH(CH ₂) ₇ CH ₃	HN NH2	O NH ₂	1665	7
314	NHCONH(CH₂)10CH3	NBoc HN NHBoc	NH ₂	1707	7
315	NHCONH(CH₂)7CH₃	HN N H	DE Z	1779	10a
316	NHCONH(CH ₂) ₇ CH ₃	HN	O NH ₂	1700	10a
317	NHCONH(CH₂)7CH₃	HN NO2	O NH2	1806	10a
318	NHCO(CH ₂) ₉ CH ₃	HN OCH ₃	O NH ₂	1793	10a
319	NHCO(CH ₂) ₉ CH ₃	HN	O NH ₂	1714	10a
320	NHCO(CH ₂) ₁₁ CH ₃	HN OCH,	O NH ₂	1821	10a
321	NHCO(CH ₂) ₁₁ CH ₃	HN NO2	ž į	1848	10a

					
322	NHCO(CH ₂) ₁₁ CH ₃	HN N	NH ₂	1742	10a
323	NHCO(CH ₂) ₈ CH ₃	HN CF3	P. S.	1943	1
324	NHCO(CH ₂) ₈ CH ₃	HN CF ₃	O NH2	2010	1
325	NHCO(CH₂)8CH₃	HN F	O NH2	1893	1
326	NHCO(CH₂) ₈ CH ₃	HN F	₹ 0= 4-	956	1
327	NHCO(CH ₂) ₈ CH ₃	HN TF	Î-()	1875	1
328	NHCO(CH ₂) ₈ CH ₃	HN CF ₃	Î	1919	1
329	NHCO(CH ₂) ₈ CH ₃	HN CF ₃	O NH ₂	1987	1
330	NHCO(CH₂)8CH₃	HN CI	Î-	1909	1
331	NHCO(CH₂)₅CH₃	HN NO ₂ CF ₃	DE Z	1998	1
332	NHCO(CH ₂) ₁₀ CH ₃	HN CCH	O H	1807	.10a
333	NHCO(CH ₂) ₁₀ CH ₃	HN NO ₂		1834	10a
334	NHCO(CH ₂) ₁₀ CH ₃	HN N	-}-	1728	10a
335	NHCONH(CH ₂) ₁₁ CH ₃	HN HN	ž į	1757	10a
336	NHCONH(CH ₂) ₁₁ CH ₃	HN NO ₂	O NH2	1864	10a
337	NHCONH(CH ₂) ₁₁ CH ₃	HN N H	O NH ₂	1836	10a

338	NHCO(CH ₂) ₁₂ CH ₃	HN NHBœ N	NH ₂	1963	10Ъ
339	NHCO(CH ₂) ₁₂ CH ₃	HN NH2 N	O NH ₂	1863	10b
340	NHCO(CH ₂) ₁₂ CH ₃	NHBo c	O NH2	2006	10ъ
341	NHCO(CH ₂) ₁₂ CH ₃	HN NH ₂	O NH2	1805	10b
342	NHCO(CH ₂) ₉ CH ₃	HN F	O NH ₂	1773	10b
343	NHCO(CH ₂) ₁₀ CH ₃	O NH ₂	O NH ₂	1786	10ь
344	NHCO(CH ₂) ₁₂ CH ₃	HN H	O NH ₂	1814	10ъ
345	NHCO(CH ₂) ₁₂ CH ₃	HN HN	O NH2	1756	10a
346	NHCO(CH ₂) ₁₂ CH ₃	HN OCH	O NH2	1836	10a
347	NHCO(CH ₂) ₇ CH ₃	HN OCH	O NH2	1765	10a
348	NHCO(CH ₂) ₇ CH ₃	HN NH	D NH2	1686	10a
349	NHCO(CH ₂) ₇ CH ₃	HN NO ₂	O NH2	1792	10a
350	HN CI	HN NHz	O NH ₂	1832	10b
351	NHCO(CH₂)11CH3	HN F	O NH ₂	1801	10jb
352	NHCONH(CH ₂) ₁₀ CH ₃	HN F	NH2	1801	10b
355	NHCONH(CH ₂) ₁₀ CH ₃	HN HN	NH ₂	1743	10a

			 		
356	NHCONH(CH ₂) ₁₀ CH ₃	HN NH OCH	NE2	1822	10a
358	NHCO(CH ₂) ₈ CH ₃	HN S H	O NH ₂	1893	1
359	NHCO(CH ₂) ₈ CH ₃	HN	O NH	948	1
360	NHCO(CH ₂) ₈ CH ₃	HN S-N NCH ₃	D NH2	938	1
361	NHCO(CH ₂) ₈ CH ₃	MN CHA	O NH ₂	952	1
362	NHCO(CH ₂) ₈ CH ₃	HN NPh	O NH ₂	969	1
363	NHCO(CH ₂) ₈ CH ₃	HM - S-N N- N	O NH ₂	970	1
364	NHCO(CH ₂) ₈ CH ₃	HM 0-3-N N-	O NH2	976	1
365	NHCO(CH₂)&CH₃	HN S-N NBn	O NH ₂	976	1
366	NHCO(CH ₂) ₈ CH ₃		₽	984	1
367	NHCO(CH₂)₃CH₃	HN Neo	O NH2	984	1
368	NHCO(CH ₂) ₈ CH ₃	0:0-N 0:0-N 0:0-N	O NH2	986	1
369	NHCO(CH ₂) ₈ CH ₃	0-b-0 0-b-0	0 NH2	987	1
370	NHCO(CH ₂) ₈ CH ₃	HZ -0-9-0	Ê-(-)	978	1
371	NHCO(CH ₂) ₈ CH ₃	HX 0:50	O N-	998	1
372	NHCO(CH ₂)₃CH₃	HN	ž	1003	1
373	NHCO(CH₂)₃CH₃	HV CI	DE 2	1003	1

374	NHCO(CH ₂) ₈ CH ₃	HM - S-N N-N	ZH.	970	1
375	NHCO(CH ₂) ₈ CH ₃	0 H - 5 O H - 5 O F	D NH	950	1
376	NHCO(CH ₂) ₈ CH ₃	HN SHA	NH ₂	950	1
377	NHCO(CH ₂) ₈ CH ₃	HN S S N F	0- - -	950	1
378	NHCO(CH ₂) ₈ CH ₃	HN	DE -	955	1
379	NHCO(CH ₂) ₈ CH ₃	HK O I	E	957	. 1
380	NHCO(CH ₂) ₈ CH ₃	HN - C	Î-	958	1
381	NHCO(CH ₂) ₈ CH ₃	0 H F	£	959	1
382	NHCO(CH ₂) ₈ CH ₃	0 H 	D= N	959	1
383	NHCO(CH ₂) ₈ CH ₃	0 HN CI	£	965	1
384	NHCO(CH ₂) ₈ CH ₃	HZ - 0 H - CI	Ē	965	1
385	NHCO(CH ₂) ₈ CH ₃	0 H 	£	975	1
386	NHCO(CH ₂) ₈ CH ₃	HN O H CF3	O NH2	975	1
387	NHCO(CH ₂) ₈ CH ₃	HN - S-N - CF3	O NH2	975	1
388	NHCO(CH ₂) ₈ CH ₃	HN 5-N F	O NH N	957	1
389	NHCO(CH ₂) ₈ CH ₃	HZ CI	£-	976	1
390	NHCO(CH ₂) ₈ CH ₃	HZ CI	\$\frac{\frac{1}{2}}{2}	976	1

- 391	NHCO(CH ₂) ₈ CH ₃	0 H CI	O NH2	976	1
392	NHCO(CH ₂) ₈ CH ₃	ocr,	O NH ₂	983	1
393	NHCO(CH ₂) ₈ CH ₃		O NH2	983	l
394	NHCO(CH ₂) ₈ CH ₃	HK - 0 - 0 - 0 - 0 - 0 - 0 - 0 - 0 - 0 -	O NH ₂	948	1
395	NHCO(CH ₂) ₈ CH ₃	HZ-0-0-0-0-0-0-0-0-0-0-0-0-0-0-0-0-0-0-0	O NH2	941	1
398	NHCO(CH₂)8CH₃	HN N F	NE.		1
399	NHCO(CH ₂) ₈ CH ₃	HN N	O NH ₂		1
400	NHCO(CH ₂) ₈ CH ₃	HN N	O NH2		1
401	NHCO(CH ₂) ₈ CH ₃	HN N N	\$ - \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \		1
402	NHCO(CH ₂) ₈ CH ₃	HN S S	\$\tilde{\text{S}}		1
403	NHCO(CH ₂) ₈ CH ₃	THE STATE OF THE S	O NH2		1
404	NHCO(CH ₂) ₈ CH ₃	HN CF ₃	NH _z		1

405	NHCO(CH ₂) ₈ CH ₃	HN \$20	O NH2	1
. 406	NHCO(CH₂)8CH₃	, H	O NH ₂	1
407	NHCO(CH ₂) ₈ CH ₃	HN CO S N CO	NH2	1
408	NHCO(CH ₂) ₈ CH ₃	HN CI	O ZE2	1
409	NHCO(CH ₂) ₈ CH ₃	HN	O NE?	. 1
410	NHCO(CH ₂) ₈ CH ₃	NH	O NH ₂	1

Preferred compounds of the present invention are compounds 45, 54, 76, 81, 85, 102, 209, 212, 253, 260, 262, 282, 285, 319, 322, 333, 334, 335, 336, 344 and 355.

According to a preferred embodiment, the present invention provides one or more crystalline forms of compounds of formula (I) and salts thereof.

Lipopeptide Intermediates

The present invention also provides compounds that are particularly useful as intermediates for the preparation of the compounds of Formula I. These compounds may also have antibacterial properties, as discussed above. In one aspect of the invention, compounds of Formula II are provided:

wherein R^{14} is selected from the group consisting of

OH
$$R^{56}$$
 and R^{56} R^{56}

wherein R^{56} is an optionally substituted straight-chain C_8 - C_{14} alkyl group and wherein q' is 0-3

In another aspect of the invention, compounds of Formula III are provided as useful intermediates for the preparation of compounds of Formula I and/or as antibacterial compounds:

wherein R¹⁵ is selected from hydrido and a carbamate amino protecting group, preferably a *tert*-butoxycarbonyl group; wherein R¹⁶ is selected from the group consisting of

wherein R^{57} is a halo or halo substituted alkyl group, preferably a fluoro or trifluoromethyl group; wherein, R^{20} is an amino acid side chain, preferably a lysine or tryptophan side chain.

Compounds 2, 10, 25, 38, 45, 50, 54, 76, 78, 79, 80, 81, 82, 84, 85, 103, 105, 107, 111, 115, 130, 138, 139, 146, 147, 150, 158, 164, 168, 174, 210, 212, 227, 253, 274, 275, 280, 283, 285, 317, 372 and 386 are useful both as antibacterial compounds and as intermediates in the synthesis of compounds of this invention.

Lipopeptide Compound Pharmaceutical Compositions and Methods of Use Thereof

Another object of the instant invention is to provide lipopeptide compounds or salts thereof, as well as pharmaceutical compositions or formulations comprising lipopeptide compounds or its salts.

Lipopeptide compounds, or pharmaceutically acceptable salts thereof, can be formulated for oral, intravenous, intramuscular, subcutaneous or parenteral administration for the therapeutic or prophylactic treatment of diseases, particularly bacterial infections. For oral or parenteral administration, lipopeptide compounds of this invention can be mixed with conventional pharmaceutical carriers and excipients and used in the form of tablets, capsules, elixirs, suspensions, syrups, wafers and the like. The compositions comprising a compound of this invention will contain from about 0.1 to about 99% by weight of the active compound, and more generally from about 10 to about 30%.

The pharmaceutical preparations disclosed herein are prepared in accordance with standard procedures and are administered at dosages that are selected to reduce, prevent or eliminate the infection (See, e. g., Remington's Pharmaceutical

Sciences, Mack Publishing Company, Easton, PA and Goodman and Gilman's The Pharmaceutical Basis of Therapeutics, Pergamon Press, New York, NY, the contents of which are incorporated herein by reference, for a general description of the methods for administering various antimicrobial agents for human therapy). The compositions of the invention (preferably of Formula I) can be delivered using controlled (e.g., capsules) or sustained release delivery systems (e.g., bioerodable matrices). Exemplary delayed release delivery systems for drug delivery that are suitable for administration of the compositions of the invention (preferably of Formula I) are described in U.S. Patent Nos. 4,452,775 (issued to Kent), 5,239,660 (issued to Leonard), 3,854,480 (issued to Zaffaroni).

The pharmaceutically-acceptable compositions of the present invention comprise one or more compounds of the invention (preferably compounds of Formula I) in association with one or more nontoxic, pharmaceutically-acceptable carriers and/or diluents and/or adjuvants and/or excipients, collectively referred to herein as "carrier" materials, and if desired other active ingredients. The compositions may contain common carriers and excipients, such as corn starch or gelatin, lactose, sucrose, microcrystalline cellulose, kaolin, mannitol, dicalcium phosphate, sodium chloride and alginic acid. The compositions may contain croscarmellose sodium, microcrystalline cellulose, corn starch, sodium starch glycolate and alginic acid.

Tablet binders that can be included are acacia, methylcellulose, sodium carboxymethylcellulose, polyvinylpyrrolidone (Povidone), hydroxypropyl methylcellulose, sucrose, starch and ethylcellulose.

Lubricants that can be used include magnesium stearate or other metallic stearates, stearic acid, silicone fluid, talc, waxes, oils and colloidal silica.

Flavoring agents such as peppermint, oil of wintergreen, cherry flavoring or the like can also be used. It may also be desirable to add a coloring agent to make the dosage form more aesthetic in appearance or to help identify the product.

For oral use, solid formulations such as tablets and capsules are particularly useful. Sustained release or enterically coated preparations may also be devised. For pediatric and geriatric applications, suspensions, syrups and chewable tablets are especially suitable. For oral administration, the pharmaceutical

compositions are in the form of, for example, a tablet, capsule, suspension or liquid. The pharmaceutical composition is preferably made in the form of a dosage unit containing a therapeutically-effective amount of the active ingredient. Examples of such dosage units are tablets and capsules. For therapeutic purposes, the tablets and capsules which can contain, in addition to the active ingredient, conventional carriers such as binding agents, for example, acacia gum, gelatin, polyvinylpyrrolidone, sorbitol, or tragacanth; fillers, for example, calcium phosphate, glycine, lactose, maize-starch, sorbitol, or sucrose, lubricants, for example, magnesium stearate, polyethylene glycol, silica, or talc; disintegrants, for example, potato starch, flavoring or coloring agents, or acceptable wetting agents. Oral liquid preparations generally are in the form of aqueous or oily solutions, suspensions, emulsions, syrups or elixirs may contain conventional additives such as suspending agents, emulsifying agents, non-aqueous agents, preservatives, coloring agents and flavoring agents. Examples of additives for liquid preparations include acacia, almond oil, ethyl alcohol, fractionated coconut oil, gelatin, glucose syrup, glycerin, hydrogenated edible fats, lecithin, methyl cellulose, methyl or propyl para-hydroxybenzoate, propylene glycol, sorbitol, or sorbic acid.

For intravenous (IV) use, a lipopeptide compound according to the invention can be dissolved or suspended in any of the commonly used intravenous fluids and administered by infusion. Intravenous fluids include, without limitation, physiological saline or Ringer's solution. Intravenous administration may be accomplished by using, without limitation, syringe, minipump or intravenous line.

Formulations for parenteral administration can be in the form of aqueous or non-aqueous isotonic sterile injection solutions or suspensions. These solutions or suspensions can be prepared from sterile powders or granules having one or more of the carriers mentioned for use in the formulations for oral administration. The compounds can be dissolved in polyethylene glycol, propylene glycol, ethanol, corn oil, benzyl alcohol, sodium chloride, and/or various buffers.

For intramuscular preparations, a sterile formulation of a lipopeptide compound or a suitable soluble salt form of the compound, for example the hydrochloride salt, can be dissolved and administered in a pharmaceutical diluent

such as Water-for-Injection (WFI), physiological saline or 5% glucose. A suitable insoluble form of the compound may be prepared and administered as a suspension in an aqueous base or a pharmaceutically acceptable oil base, e.g., an ester of a long chain fatty acid such as ethyl oleate.

A dose of an intravenous, intramuscular or parental formulation of a lipopeptide compound may be adminstered as a bolus or by slow infusion. A bolus is a dose that is administered in less than 30 minutes. In a preferred embodiment, a bolus is administered in less than 15 or less than 10 minutes. In a more preferred embodiment, a bolus is administered in less than 5 minutes. In an even more preferred embodiment, a bolus is administered in one minute or less. An infusion is a dose that is administered at a rate of 30 minutes or greater. In a preferred embodiment, the infusion is one hour or greater. In another embodiment, the infusion is substantially constant.

For topical use the compounds of the present invention can also be prepared in suitable forms to be applied to the skin, or mucus membranes of the nose and throat, and can take the form of creams, ointments, liquid sprays or inhalants, lozenges, or throat paints. Such topical formulations further can include chemical compounds such as dimethylsulfoxide (DMSO) to facilitate surface penetration of the active ingredient.

For application to the eyes or ears, the compounds of the present invention can be presented in liquid or semi-liquid form formulated in hydrophobic or hydrophilic bases as ointments, creams, lotions, paints or powders.

For rectal administration the compounds of the present invention can be administered in the form of suppositories admixed with conventional carriers such as cocoa butter, wax or other glyceride.

Alternatively, the compounds of the present invention can be in powder form for reconstitution in the appropriate pharmaceutically acceptable carrier at the time of delivery. In another embodiment, the unit dosage form of the compound can be a solution of the compound or preferably a salt thereof in a suitable diluent in sterile, hermetically sealed ampoules or sterile syringes. The concentration of the compound in the unit dosage may vary, e.g. from about 1 percent to about 50

percent, depending on the compound used and its solubility and the dose desired by the physician. If the compositions contain dosage units, each dosage unit preferably contains from 1-500 mg of the active material. For adult human treatment, the dosage employed preferably ranges from 5 mg to 10 g, per day, depending on the route and frequency of administration.

In another aspect, the invention provides a method for inhibiting the growth of microorganisms, preferably bacteria, comprising contacting said organisms with a compound of the invention, preferably a compound of Formula I, under conditions which permit entry of the compound into said organism and into said microorganism. Such conditions are known to one skilled in the art and are exemplified in the Examples. This method involves contacting a microbial cell with a therapeutically-effective amount of compound(s) of the invention, preferably compound(s) of Formula I, in vivo or in vitro.

According to this aspect of the invention, the novel compositions disclosed herein are placed in a pharmaceutically acceptable carrier and are delivered to a recipient subject (preferably a human) in accordance with known methods of drug delivery. In general, the methods of the invention for delivering the compositions of the invention *in vivo* utilize art-recognized protocols for delivering the agent with the only substantial procedural modification being the substitution of the compounds of the invention (preferably compounds of Formula I) for the drugs in the art-recognized protocols. Likewise, the methods for using the claimed composition for treating cells in culture, for example, to eliminate or reduce the level of bacterial contamination of a cell culture, utilize art-recognized protocols for treating cell cultures with antibacterial agent(s) with the only substantial procedural modification being the substitution of the compounds of the invention (preferably compounds of Formula I) for the agents used in the art-recognized protocols.

In one embodiment, the invention provides a method for treating an infection, especially those caused by gram-positive bacteria, in a subject with a therapeutically-effective amount of a lipopeptide compound according to Formula I. Exemplary procedures for delivering an antibacterial agent are described in U.S. Patent No. 5,041,567, issued to Rogers and in PCT patent application number

EP94/02552 (publication no. WO 95/05384), the entire contents of which documents are incorporated in their entirety herein by reference. As used herein the phrase "therapeutically-effective amount" means an amount of a compound of the present invention that prevents the onset, alleviates the symptoms, or stops the progression of a bacterial infection. The term "treating" is defined as administering, to a subject, a therapeutically-effective amount of a compound of the invention (preferably a compound of Formula I) both to prevent the occurrence of an infection and to control or eliminate an infection. The term "subject", as described herein, is defined as a mammal, a plant or a cell culture. In a preferred embodiment, a subject is a human or other animal patient in need of lipopeptide compound treatment.

The method comprises administering to the subject an effective dose of a compound of this invention. An effective dose is generally between about 0.1 and about 100 mg/kg of a lipopeptide compound of Formula I or a pharmaceutically acceptable salt thereof. A preferred dose is from about 0.1 to about 50 mg/kg of a lipopeptide compound of Formula I or a pharmaceutically acceptable salt thereof. A more preferred dose is from about 1 to 25 mg/kg of a lipopeptide compound of Formula I or a pharmaceutically acceptable salt thereof. An effective dose for cell culture is usually between 0.1 and 1000 μ g/mL, more preferably between 0.1 and 200 μ g/mL.

The compound of Formula I can be administered as a single daily dose or in multiple doses per day. The treatment regime may require administration over extended periods of time, e.g., for several days or for from two to four weeks. The amount per administered dose or the total amount administered will depend on such factors as the nature and severity of the infection, the age and general health of the patient, the tolerance of the patient to the compound and the microorganism or microorganisms involved in the infection. A method of administration to a patient of daptomycin, another member of the lipopeptide compound class, is disclosed in United States Serial No. 09/406,568, filed September 24, 1999, which claims the benefit of U.S. Provisional Application Nos. 60/101,828, filed September 25, 1998, and 60/125,750, filed March 24, 1999.

A lipopeptide compound according to this invention may also be administered in the diet or feed of a patient or animal. If administered as part of a total dietary intake, the amount of compound employed can be less than 1% by weight of the diet and preferably no more than 0.5% by weight. The diet for animals can be normal foodstuffs to which the compound can be added or it can be added to a premix.

The methods of the present invention comprise administering a lipopeptide compound of Formula I or a pharmaceutical composition thereof to a subject in need thereof in an amount that is efficacious in reducing or eliminating the bacterial infection. The compound may be administered orally, parenterally, by inhalation, topically, rectally, nasally, buccally, vaginally, or by an implanted reservoir, external pump or catheter. The compound may be prepared for opthalmic or aerosolized uses. The compounds of the present invention can be administered as an aerosol for the treatment of pneumonia or other lung-based infections. A preferred aerosol delivery vehicle is an anhydrous or dry powder inhaler. Lipopeptide compounds of Formula I or a pharmaceutical composition thereof also may be directly injected or administered into an abscess, ventricle or joint. Parenteral administration includes subcutaneous, intravenous, intramuscular, intra-articular, intra-synovial, cisternal, intrathecal, intrahepatic, intralesional and intracranial. injection or infusion. In a preferred embodiment, lipopeptide compounds are administered intravenously, subcutaneously or orally. In a preferred embodiment for administering a lipopeptide compound according to Formula I to a cell culture, the compound may be administered in a nutrient medium.

The method of the instant invention may be used to treat a subject having a bacterial infection in which the infection is caused or exacerbated by any type of bacteria, particularly gram-positive bacteria. In one embodiment, a lipopeptide compound or a pharmaceutical composition thereof is administered to a patient according to the methods of this invention. In a preferred embodiment, the bacterial infection may be caused or exacerbated by gram-positive bacteria. These gram-positive bacteria include, but are not limited to, methicillin-susceptible and methicillin-resistant staphylococci (including *Staphylococcus aureus*, *S. epidermidis*,

S. haemolyticus, S. hominis, S. saprophyticus, and coagulase-negative staphylococci), glycopeptide intermediary- susceptible S. aureus (GISA), penicillin-susceptible and penicillin-resistant streptococci (including Streptococcus pneumoniae, S. pyogenes, S. agalactiae, S. avium, S. bovis, S. lactis, S. sangius and Streptococci Group C, Streptococci Group G and viridans streptococci), enterococci (including vancomycin-susceptible and vancomycin-resistant strains such as Enterococcus faecalis and E. faecium), Clostridium difficile, C. clostridiiforme, C. innocuum, C. perfringens, C. ramosum, Haemophilus influenzae, Listeria monocytogenes, Corynebacterium jeikeium, Bifidobacterium spp., Eubacterium aerofaciens, E. lentum, Lactobacillus acidophilus, L. casei, L. plantarum, Lactococcus spp., Leuconostoc spp., Pediococcus, Peptostreptococcus anaerobius, P. asaccarolyticus, P. magnus, P. micros, P. prevotii, P. productus, Propionibacterium acnes, Actinomyces spp., Moraxella spp. (including M. catarrhalis) and Escherichia spp. (including E. coli).

In a preferred embodiment, the antibacterial activity of lipopeptide compounds of Formula I against classically "resistant" strains is comparable to that against classically "susceptible" strains in *in vitro* experiments. In another preferred embodiment, the minimum inhibitory concentration (MIC) value for lipopeptide compounds according to this invention against susceptible strains is typically the same or lower than that of vancomycin. Thus, in a preferred embodiment, a lipopeptide compound of this invention or a pharmaceutical composition thereof is administered according to the methods of this invention to a patient who exhibits a bacterial infection that is resistant to other compounds, including vancomycin or daptomycin. In addition, unlike glycopeptide antibiotics, lipopeptide compounds exhibits rapid, concentration-dependent bactericidal activity against gram-positive organisms. Thus, in a preferred embodiment, a lipopeptide compound according to this invention or a pharmaceutical composition thereof is administered according to the methods of this invention to a patient in need of rapidly acting antibiotic therapy.

The method of the instant invention may be used for any bacterial infection of any organ or tissue in the body. In a preferred embodiment, the bacterial infection is caused by gram-positive bacteria. These organs or tissue include, without limitation, skeletal muscle, skin, bloodstream, kidneys, heart, lung and bone. The

method of the invention may be used to treat, without limitation, skin and soft tissue infections, bacteremia and urinary tract infections. The method of the invention may be used to treat community acquired respiratory infections, including, without limitation, otitis media, sinusitis, chronic bronchitis and pneumonia, including pneumonia caused by drug-resistant *S. pneumoniae* or *H. influenzae*. The method of the invention also may be used to treat mixed infections that comprise different types of gram-positive bacteria, or which comprise both gram-positive and gram-negative bacteria. These types of infections include intra-abdominal infections and obstetrical/gynecological infections. The method of the invention also may be used to treat an infection including, without limitation, endocarditis, nephritis, septic arthritis, intra-abdominal sepsis, bone and joint infections. and osteomyelitis. In a preferred embodiment, any of the above-described diseases may be treated using lipopeptide compounds according to this invention or pharmaceutical compositions thereof.

The method of the instant invention may also be practiced while concurrently administering one or more other antimicrobial agents, such as antibacterial agents (antibiotics) or antifungal agents. In one aspect, the method may be practiced by administering more than one lipopeptide compounds according to this invention. In another embodiment, the method may be practiced by administering a lipopeptide compound according to this invention with another lipopeptide compound, such as daptomycin.

Antibacterial agents and classes thereof that may be co-administered with a compound of the present invention include, without limitation, penicillins and related drugs, carbapenems, cephalosporins and related drugs, aminoglycosides, bacitracin, gramicidin, mupirocin, chloramphenicol, thiamphenicol, fusidate sodium, lincomycin, clindamycin, macrolides, novobiocin, polymyxins, rifamycins, spectinomycin, tetracyclines, vancomycin, teicoplanin, streptogramins, anti-folate agents including sulfonamides, trimethoprim and its combinations and pyrimethamine, synthetic antibacterials including nitrofurans, methenamine mandelate and methenamine hippurate, nitroimidazoles, quinolones, fluoroquinolones, isoniazid, ethambutol, pyrazinamide, para-aminosalicylic acid (PAS), cycloserine, capreomycin, ethionamide, prothionamide, thiacetazone,

viomycin, eveminomycin, glycopeptide, glycylcylcline, ketolides, oxazolidinone; imipenen, amikacin, netilmicin, fosfomycin, gentamicin, ceftriaxone, Ziracin, LY 333328, CL 331002, HMR 3647, Linezolid, Synercid, Aztreonam, and Metronidazole, Epiroprim, OCA-983, GV-143253, Sanfetrinem sodium, CS-834, Biapenem, A-99058.1, A-165600, A-179796, KA 159, Dynemicin A, DX8739, DU 6681; Cefluprenam, ER 35786, Cefoselis, Sanfetrinem celexetil, HGP-31, Cefpirome, HMR-3647, RU-59863, Mersacidin, KP 736, Rifalazil; Kosan, AM 1732, MEN 10700, Lenapenem, BO 2502A, NE-1530, PR 39, K130, OPC 20000, OPC 2045, Veneprim, PD 138312, PD 140248, CP 111905, Sulopenem, ritipenam acoxyl, RO-65-5788, Cyclothialidine, Sch-40832, SEP-132613, micacocidin A, SB-275833, SR-15402, SUN A0026, TOC 39, carumonam, Cefozopran, Cefetamet pivoxil, and T 3811.

In a preferred embodiment, antibacterial agents that may be coadministered with a compound according to this invention include, without limitation, imipenen, amikacin, netilmicin, fosfomycin, gentamicin, ceftriaxone, teicoplanin, Ziracin, LY 333328, CL 331002, HMR 3647, Linezolid, Synercid, Aztreonam, and Metronidazole.

Antifungal agents that may be co-administered with a compound according to this invention include, without limitation, Caspofungen, Voriconazole, Sertaconazole, IB-367, FK-463, LY-303366, Sch-56592, Sitafloxacin, DB-289 polyenes, such as Amphotericin, Nystatin, Primaricin; azoles, such as Fluconazole, Itraconazole, and Ketoconazole; allylamines, such as Naftifine and Terbinafine; and anti-metabolites such as Flucytosine. Other antifungal agents include without limitation, those disclosed in Fostel et al., Drug Discovery Today 5:25-32 (2000), herein incorporated by reference. Fostel et al. disclose antifungal compounds including Corynecandin, Mer-WF3010, Fusacandins, Artrichitin/LL 15G256γ, Sordarins, Cispentacin, Azoxybacillin, Aureobasidin and Khafrefungin.

Lipopeptide compounds may be administered according to this method until the bacterial infection is eradicated or reduced. In one embodiment, a lipopeptide compound is administered for a period of time from 3 days to 6 months. In a preferred embodiment, a lipopeptide compound is administered for 7 to 56 days.

In a more preferred embodiment, a lipopeptide compound is administered for 7 to 28 days. In an even more preferred embodiment, a lipopeptide compound is administered for 7 to 14 days. Lipopeptide compounds may be administered for a longer or shorter time period if it is so desired.

General Procedures for Lipopeptide Compound Synthesis

Lipopeptide compounds of Formula I may be produced as described below. The lipopeptide compounds of the instant invention may be produced semi-synthetically using daptomycin as a starting point or may be produced by a total synthesis approach.

For the semi-synthetic approach according to the present invention, daptomycin may be prepared by any method known in the art. See, e.g., United States Patents 4,885,243 and 4,874,843. Daptomycin may be used in its acylated state or it may be deacylated prior to its use as described herein. Daptomycin may be deacylated using *Actinoplanes utahensis* as described in United States Patent 4,482,487. Alternatively, daptomycin may be deacylated as follows:

Daptomycin (5.0 g) was dissolved in water (25 ml) and adjusted to pH 9 with 5M sodium hydroxide. Ditert-butyldicarbonate (1.5 g) was added and the mixture was adjusted to maintain pH 9 with 5 M sodium hydroxide until the reaction was complete (4 hours). The pH was adjusted to 7 and the mixture was loaded onto a Bondesil 40µ C8 resin column. The column was washed with water and the product was eluted from the column with methanol. Evaporation of the methanol gave BOC-protected daptomycin as a yellow powder.

A preparation of deacylase enzyme was produced from recombinant Streptomyces lividans, which expresses the Actinoplanes utahensis deacylase enzyme. The enzyme in ethylene glycol (400 µl) was added to BOC-protected daptomycin (1 g) in water (100 ml) at pH 7-8. After incubation for 72 hours, the mixture was loaded on a Bondesil 40µ C8 resin column. The column was washed with water and the product was eluted from the column with 10% acetonitrile in water. The product was evaporated to give deacylated BOC-protected daptomycin as a yellow powder.

Kynurenine Derivatives

Scheme 1

Daptomycin can be converted into analogs bearing modifications at the R² position by converting the aromatic amino group to the diazonium salt compound I with reagents such as sodium nitrite/hydrochloric acid or isoamylnitrite. Using chemistry known to those skilled in the art and following the teachings of the disclosure, the diazonium group can then be displaced by reagents such as sodium azide, potassium ethylxanthate or copper chloride to yield derivative compounds II, wherein R¹⁹ is as previously defined.

Scheme 2

Additionally, compound I can be converted to the azide compound III by reaction with an azide source, typically sodium azide. Modifications to the ketone group can then be undertaken using chemistry known to those having ordinary skill in the art, such as reduction, oxime formation, ketalization conversion to a leaving group and displacement to give compounds of formula IV, wherein R¹⁷ and R¹⁸ are as previously defined.

Scheme 3

Compound IV may also be converted to compound V by reducing the azide group to the amine using chemistry known to those having ordinary skill in the art, and following the teachings of the disclosure, such as reaction with triphenyl phosphine and water, or reducing agents such as sodium borohydride wherein R^{17} and R^{18} are as previously defined.

Additionally compound I can be converted into compound VI by reduction with hypophosphorus acid. Modifications to the ketone group can then be undertaken using chemistry known to those having ordinary skill in the art similar to those used in scheme 2, wherein R¹⁷ and R¹⁸ are as previously defined.

Ornithine derivatives

Scheme 1

Daptomycin can be converted into analogs bearing modifications at the R¹ position by treating the aromatic amino group of the ornithine with reagents such as isocyanates, isothiocyanates, activated esters, acid chlorides, sulfonylchlorides or activated sulfonamides, heterocycles bearing readily displaceable groups, imidates, lactones or reductively with aldehydes to yield compound VIII, wherein R¹ is as previously defined.

Tryptophan Amine Derivatives

Scheme 1

Daptomycin can be converted into compound IX by first protecting the ornithine amine with an appropriate amino protecting group (P) known to those skilled in the art and following the teachings of the disclosure. The decyl side chain on the tryptophan is then removed using an enzyme capable of deacylating daptomycin, such as that described above.

Scheme 2

Compound IX can be modified at the tryptophan amine with reagents such as isocyanates, isothiocyanates, activated esters, acid chlorides, sulfonylchlorides or activated sulfonamides, heterocycles bearing readily displaceable groups, imidates, lactones or reductively with aldehydes to yield compound X. Compound X can be deprotected to give compound XI according to procedures known to those skilled in the art following the disclosure of this invention, wherein R is as previously defined.

The above modifications to the ornithine amine R¹, tryptophan amine R or kynurenine side chain R² may be independently combined to yield additional compounds that are modified at up to all three sites. In order to achieve these modifications, it may be necessary to protect certain functionalities in the molecule. Protecting these functionalities should be within the expertise of one skilled in the art following the disclosure of this invention. See, e.g., Greene, *supra*.

Solid Support Synthesis of Lipopeptide Compounds

In an alternative embodiment of the invention, the lipopeptide compounds of Formula I may be synthesized on a solid support as outlined below. In step 1, a suitably-N-protected-βMeGlu(OH)-OAllyl ester is coupled to a suitable resin to give Compound XII. Deprotection of the amino group of Compound XII, followed by coupling of the amino group with a suitably protected seryl derivative (A1) gives Compound XIII, wherein P is a suitable protecting group. This peptide coupling process, i.e., deprotection of the alpha-amino group, followed by coupling to a suitably protected amino acid, is repeated until the desired number of amino acids have been coupled to the resin. In the scheme shown below, eleven amino acids have been coupled to give Compound XIV. Addition of an activated R group, R*, is added to Compound XIV to give Compound XV. In step 4, Compound XV is cyclized to give Compound XVI. Subsequently, in step 5, Compound XVI is removed from the resin to give the lipopeptide Compound XVII.

Synthetic Scheme for Total Synthesis of Lipopeptide Compounds

$$A^{I} = \bigvee_{HN \setminus I}^{31} O$$

, wherein A^1 , is a suitably protected serine derivative, wherein R^{31} is a suitable, cleavable hydroxyl protecting group as outlined below.

$$A^2 = A^7 =$$

, wherein A² and A⁷, are suitably protected glycine derivatives as

outlined below.

$$A^{3} = A^{5} = A^{5$$

protected aspartic acid derivatives as outlined below, wherein ²⁸R, ²⁹R and ³⁰R are cleavable protecting groups, preferably t-butyl groups.

$$A^4 = \bigvee_{HN, \uparrow} O$$

, wherein A⁴ is a suitably protected alanine derivative as outlined

below.

$$A^6 = \begin{matrix} *R^1 & \checkmark \\ HN_{\sim} & \checkmark \\ HN_{\sim} & \checkmark \end{matrix}$$

wherein A^6 is a suitably protected ornithine derivative as outlined below, or derivatized ornthine wherein $*R^1$ is R^1 as previously described or alternatively a protected form of R^1 that would yield R^1 upon subsequent deprotection.

, wherein A⁸ is a suitably protected depsipeptide as outlined below, Y is a protecting group that is cleavable under conditions that leave other protecting groups intact to the others used, i.e., Alloc, and wherein *R² is R² as previously described or alternatively a protected form of R² that would yield R² upon subsequent deprotection. Preferably ²*R is a kynurenine, or substituted kynurenine side chain, most preferably

$$R^2 = 0$$
 NH_2

$$A^{10} = H_2N O O$$

wherein A¹⁰ is a suitably protected asparagine derivative as

outlined below.

$$A^{11} = \bigvee_{HNC}^{N} R^{37}$$

wherein A^{11} is a suitably protected tryptophan derivative as outlined below, wherein R^{*37} is hydrido or a suitable protecting group, preferably t-butoxy carbonyl.

It will be understood by those skilled in the art that both the amino and the side chain functional groups must be suitably protected prior to attaching them to the growing peptide chain. Suitable protecting groups can be any group known in the art to be useful in peptide synthesis. Such pairings of protecting groups are well known. See, e.g., "Synthesis Notes" in the Novabiochem Catalog and Peptide Synthesis Handbook (1999), pages S1-S93 and references cited therein. Following the disclosure of the present application, the selection of protecting groups and method of use thereof will be known to one skilled in the art.

It will also be understood by those skilled in the art that the choice of protecting group on the side chain functional groups will either result or not result in the protecting group being cleaved concomitantly with the peptide's final cleavage from the resin, which will give the natural amino acid functionality or a protected derivative thereof, respectively.

The following general procedures serve to exemplify the solid support synthesis of compounds of Formula I.

Step 1: Coupling suitably-N-protected-\(\beta MeGlu(OH)\)-OAllyl ester to a resin

Five molar equivalents each, with respect to the resin, of a suitably-N-protected-βMeGlu(OH)-OAllyl ester, 1,3-Diisopropylcarbodiimide (DIC) and 1-Hydroxy-7-azabenzotriazole (HOAt) are stirred for 30 mins in dimethylformamide (DMF; 5ml/g resin). A suitably functionalised resin or solid support, such as, but not limited to, Wang, Safety Catch, Rink, Knorr, PAL, or PAM resin, is added and the resulting suspension is stirred for 16 hrs. The resin-N-protected-βMeGlu(OH)-OAllyl ester is then filtered, dried and the coupling is repeated. The N-protecting group is then removed using the appropriate conditions given in the coupling steps below.

Step 2: (A) General coupling cycle for amino acids with an N-9-Fluorenylmethoxycarbonyl (Fmoc) protecting group

Five molar equivalents each, with respect to the resin-AA(wherein resin-AA is defined as the resin attached the the growing amino acid chain), of a suitably protected Fmoc amino acid, DIC, and HOAt (0.5 molar solution in DMF) are added to the resin-AA, along with sufficient DMF to give a working volume. The mixture is shaken for one hour, filtered, and the coupling is repeated. After the second coupling the resin is washed twice with DMF, twice with methanol, and twice again with DMF. The Fmoc group of the newly coupled amino acid A¹⁻¹¹ is deprotected by stirring the resin product in one working volume of a solution of 20% piperidine in N-methyl pyrolidine for five minutes, filtering the resin, and stirring the resin in 20% piperidine in N-methyl pyrolidine again for 20 minutes. The resin is washed twice with DMF, twice with methanol, and twice again with DMF.

Step 2 (B): General coupling cycle of amino acids with an N-tert-Butoxy-carbonyl (N-Boc) protecting group

Five molar equivalents each, with respect to the resin-AA, of a suitably protected N-Boc amino acid, DIC, and HOAt (0.5 molar solution in DMF) are added to the resin-AA, along with sufficient DMF to give a working volume. The mixture is shaken for one hour, filtered, and the coupling is repeated. After the repeated coupling the resin is washed twice with DMF, twice with methanol, and twice again

with DMF. The Boc group of the newly coupled amino acid A¹⁻¹¹, is then deprotected by stirring the resin in one working volume of CH₂Cl₂:trifluoroacetic acid (TFA) 1:1 for 15 minutes, filtering, and stirring in one working volume of CH₂Cl₂:TFA 1:1 for another 15 minutes. The resin is neutralized by washing with excess diisopropylethylamine (DIPEA) in CH₂Cl₂ and then washed twice with DMF, twice with methanol, and twice again with DMF.

Step 3: Terminal amine capping reaction

Ten molar equivalents, with respect to the resin XV, of a suitable reagent containing R* such as an activated ester, isocyanate, thioisocyanate, anhydride, acid chloride, chloroformate, or reactive salt thereof, in one working volume of DMF is added to the resin XIV and agitated for 25 hours. The resulting resin XV is washed twice with DMF, twice with methanol, and twice again with DMF.

Step 4: Cyclization

The dried resin XV is placed under an argon atmosphere, and treated with a solution of Pd(PPh₃)₄ 125 mgs/0.1 mmol peptide substrate, in CH₂Cl₂: Acetic acid: N-Methylmorpholine, 40: 2: 1, 1 ml/0.1 mmol peptide substrate. The mixture is stirred for 3 hours at ambient temperature, filtered, and washed twice with DMF, twice with methanol, and twice again with DMF. Five molar equivalents each, with respect to the resin, of DIC, and HOAt (0.5 molar solution in DMF) are added to the resin, along with sufficient DMF to give a working volume. The reaction is shaken for 17 hours, filtered, and washed twice with DMF, twice with methanol, and twice again with DMF to give resin XVI.

Step 5: Cleavage and isolation of the lipopeptide

The desired lipopeptide is cleaved from resin XVI and isolated, resulting in a compound in which R^{27} is OH or NH_2 . If Fmoc chemistry is used, the dried resin is suspended in 1 ml / 0.1 mmol peptide substrate of CH_2Cl_2 : TFA: Ethanedithiol (EDT): Triisopropylsilane (TIS), 16:22:1:1, and stirred for 6-8

hours at ambient temperature. The resin is filtered, washed with 1 equal volume of cold TFA, and the combined filtrates are evaporated under reduced pressure. Crude product XVII is then precipitated by the addition of diethyl ether, and isolated by centrifugation. This product may be further purified by preparative reverse phase HPLC.

If N-Boc chemistry is used, the dried resin is suspended in hydrogen flouride (HF): anisole: dimethylsulfide (DMS), 10:1:1, and stirred for 2 hours at 0° C. The volitiles are evaporated under a stream of nitrogen. The resin is then extracted with TFA, filtered and washed twice with TFA, and the combined TFA filtrates evaporated under reduced pressure. Crude product is then precipitated by the addition of diethyl ether, and isolated by centrifugation. This product may be further purified by preparative reverse phase HPLC.

If the resin is a Safety Catch resin, then $R^{27} = OR$ or NRH. The dried resin XVI is suspended in N-methylpyrolidine (NMP) or dimethylsulphoxide (DMSO) (8 ml/g resin), Five equivalents of DIPEA (with respect to resin substitution) and 24 equivalents of iodo or bromoacetonitrile (with respect to resin substitution) are added. The suspension is stirred for 24 hours at ambient temperature under inert atmosphere. The resin is filtered, washed with tetrahydrofuran (THF) and DMSO. For an ester, the resin is then treated with an alcohol, hydroxide or alkoxide (20 equivalents with respect to resin substitution) in THF for 20 hours. The resin is filtered, washed with THF and water, and the combined filtrates are evaporated under reduced pressure. Crude product is precipitated by the addition of diethyl ether, and isolated by centrifugation. The product may be further purified by preparative reverse phase HPLC. For amides the resin is then treated with a primary or secondary amine (20 equivalents with respect to resin substitution) in THF for 12-40 hours, at a gentle reflux under inert atmosphere. The resin is filtered, washed with THF and water, and the combined filtrates are evaporated under reduced pressure. Crude product is then precipitated by the addition of diethyl ether, and isolated by centrifugation. This product may be further purified by preparative reverse phase HPLC.

In order that this invention may be more fully understood, the following examples are set forth. These examples are for the purpose of illustration only and are not to be construed as limiting the scope of the invention in any way.

EXAMPLE 1 - PREPARATION OF COMPOUNDS 38, 40, 50, 52, 77-80, 82-84, 87-100, 103-169, 171-176, 183-187, 194-199, 201-204, 208, 210-211, 222-244, 252, 265-267, 271-281, 283-284, 286-291, 323-331, 358-395 and 398-410

A suspension of daptomycin in dry dimethylformamide (0.6 ml) was treated with a solution of 4-Fluorobenzaldehyde (0.2 ml) and a suspension of sodium triacetoxyborohydride (0.2 ml, 1.5M in dry dimethylformamide). After 24 hours, the reaction mixture was diluted with water/acetonitrile (1:1; 0.4 ml) and purified by preparative HPLC. The reaction mixture was loaded onto an IBSIL-C8 5μ 250x20.2mm column and eluted at 20 ml/min with 30-60% acetonitrile in 5mM ammonium phosphate buffer. Fractions containing product were collected and freezedried. The freeze-dried residue was dissolved in water (5 ml) and applied to a Bondesil 40μ C8 resin column. The column was washed with water and eluted with methanol. Evaporation of the methanol gave compound 38 as a pale yellow solid (23 mg).

In an analogous manner, compounds 40, 50, 52, 77-80, 82-84, 87-100, 103-169, 171-176, 183-187, 194-199, 201-204, 208, 210-211, 222-244, 252, 265-267, 271-281, 283-284, 286-291, 323-331, 358-395 and 398-410 can be prepared as detailed in the above example by appropriate substitutions of reagents obvious to those skilled in the art following the teachings of the disclosure.

EXAMPLE 1a - PREPARATION OF COMPOUND 282

2-Methyl-6-nitroquinoline (0.4ml, 0.5M solution in dioxane) was treated with selenium dioxide (0.2 ml, 0.9M solution in 9/1 dioxane/water) and heated to 90°C overnight. The mixture was cooled to room temperature and diluted with water (1 ml). The mixture was then extracted with ethyl acetate (3 x 2 ml). The organic extract was then dried in vacuo to give 6-nitro-2-quinolinecarboxaldehyde which was carried forward without further purification. Daptomycin (1 ml, 0.1 M in

dry dimethylformamide) was treated successively with 6-nitro-2-quinolinecarboxaldehyde prepared above in dry dimethylformamide (0.2 ml) and sodium triacetoxyborohydride (0.4 ml, 1.5M solution in dry dimethylformamide). The mixture was capped and shaken briefly. After 24h, the mixture was treated with water (0.2 ml) and loaded onto an IBSIL-C8 5μ 250 x 20.2mm column. The column was eluted at 25ml/min under the gradient conditions of 30-60% acetonitrile in 5mM ammonium phosphate buffer over 25 min followed by holding at 60% acetonitrile in 5mM ammonium phosphate buffer for another 10 min. The desired fractions were collected and the acetonitrile was removed by evaporation. The residue was applied to a Bondesil 40μ C8 resin column, washed with water and the product was eluted with methanol. Evaporation of the methanol gave compound 282 as a pale yellow solid.

EXAMPLE 1b - PREPARATION OF COMPOUND 285

4-Chloro-2-methylquinoline (0.4ml, 0.5M solution in dioxane) was treated with selenium dioxide (0.2 ml, 0.9M solution in 9/1 dioxane/water) and heated to 90°C overnight. The mixture was cooled to room temperature and diluted with water (1 ml). The mixture was then extracted with ethyl acetate (3 x 2 ml). The organic extract was then dried in vacuo to give 4-chloro-2-quinolinecarboxaldehyde which was carried forward without further purification. Daptomycin (1ml, 0.1 M in dry dimethylformamide) was treated successively with 4-chloro-2quinolinecarboxaldehyde prepared above and diluted in dry dimethylformamide (0.2 ml) and sodium triacetoxyborohydride (0.4 ml, 1.5M in dry dimethylformamide). The mixture was capped and shaken briefly. After 24h the mixture was treated with water (0.2 ml) and loaded on an IBSIL-C8 5μ 250 x 20.2mm column. The column was eluted at 25ml/min under the gradient conditions of 30-60% acetonitrile in 5mM ammonium phosphate buffer over 25 min followed by holding at 60% acetonitrile in 5mM ammonium phosphate buffer for another 10 min. The desired fractions were collected and the acetonitrile was removed by evaporation. The residue was applied to a Bondesil 40μ C8 resin column, washed with water and the product eluted off with methanol. Evaporation of the methanol gave compound 285 as a yellow solid.

EXAMPLE 1c - PREPARATION OF COMPOUND 85

Daptomycin (1ml, 0.1M in dry dimethylformamide) was treated successively with 1-methyl-2-imidazolecarboxaldehyde (0.2 ml, 0.5M solution in dry dimethylformamide) and sodium triacetoxyborohydride (0.4 ml, 1.5M solution in dry dimethylformamide). The mixture was capped and shaken briefly. After 24h the mixture was treated with water (0.2 ml) and loaded onto an IBSIL-C8 5µ 250 x 20.2mm column. The column was eluted at 30 ml/min under the gradient conditions of 35-40% acetonitrile in 5mM ammonium phosphate buffer over 30 min. The desired fractions were collected and the acetonitrile was removed by evaporation. The residue was applied to a Bondesil 40µ C8 resin column, washed with water and eluted with methanol. This mixture was then loaded on a Prodigy ODS 10µ 250 x 21.2mm column eluted at 50 ml/min at 33% acetonitrile in 5mM ammonium phosphate buffer adjusted to pH 3.2. The desired fractions were collected and the acetonitrile was removed by evaporation. The residue was applied to a Bondesil 40µ C8 resin column, washed with water and the product was eluted with methanol. Evaporation of the methanol gave compound 85 as a pale yellow solid.

EXAMPLE 1d - PREPARATION OF COMPOUND 212

Daptomycin (1ml, 0.1 M in dry dimethylformamide) was treated successively with 2-imidazolecarboxaldehyde (0.2 ml, 0.5M solution in dry dimethylformamide) and sodium triacetoxyborohydride (0.4 ml, 1.5M solution in dry dimethylformamide). The mixture was capped and shaken briefly After 24h, the mixture was treated with water (0.2 ml) and the mixture was loaded on an IBSIL-C8 5μ 250 x 20.2mm column. The column was eluted at 30 ml/min under the gradient conditions of 35-40% acetonitrile in 5mM ammonium phosphate buffer over 30 min. The desired fractions were collected and the acetonitrile was removed by evaporation. The residue was applied to a Bondesil 40μ C8 resin column, washed with water and eluted with methanol. This mixture was then loaded on a Prodigy ODS 10μ 250 x 21.2mm column and eluted at 50 ml/min at 33% acetonitrile in 5mM ammonium phosphate buffer adjusted to pH 3.2. The desired fractions were collected and the acetonitrile was removed by evaporation. The residue was applied to a Bondesil 40μ

C8 resin column, washed with water and the product eluted with methanol. Evaporation of the methanol gave compound 212 as a yellow solid.

EXAMPLE 1e - PREPARATION OF COMPOUND 81

Daptomycin (1ml, 0.1M in dry dimethylformamide) was treated successively with 5-fluoroindole-3-carboxaldehyde (0.2 ml, 0.5M solution in dry dimethylformamide) and sodum triacetoxyborohydride (0.4 ml, 1.5M solution in dry dimethylformamide). The mixture was capped and shaken briefly. After 24h the mixture was treated with water (0.2 ml) and loaded on an IBSIL-C8 5μ 250 x 20.2mm column. The column was eluted at 25 ml/min under the gradient conditions of 30-60% acetonitrile in 5mM ammonium phosphate buffer over 25 min followed by holding at 60% acetonitrile in 5mM ammonium phosphate buffer for another 10 min. The desired fractions were collected, the acetonitrile was removed by evaporation and the residue applied to a Bondesil 40μ C8 resin column. The column was washed with water and the product was eluted with methanol. Evaporation of the methanol gave compound 81 as a pale yellow solid.

EXAMPLE If - PREPARATION OF COMPOUND 253

p-N,N-Bis(2-chloroethyl)aminobenzaldehyde (0.3g) was dissolved in acetone (2.5 ml) and treated with sodium iodide (0.4g). The mixture was warmed to 40°C for 3h then treated with benzylamine (0.2 ml) and triethylamine (0.4 ml). The mixture was diluted to 7 ml with acetonitrile and then heated to 60°C. After 24h, the mixture was cooled to room temperature and the solvent was removed by evaporation. 4-(4-Benzylpiperazino)benzaldehyde was purified by silica gel chromatography eluting with (10% triethylamine/methanol/dichloromethane).

Daptomycin (1ml, 0.1 M in dry dimethylformamide) was treated successively with the 4-(4-benzylpiperazino)benzaldehyde prepared above diluted in dry dimethylformamide (0.2 ml), and sodium triacetoxyborohydride (0.4 ml, 1.5M solution in dry dimethylformamide). The mixture was capped and shaken briefly. After 24h the mixture was treated with water (0.2 ml) and loaded on an IBSIL-C8 5μ 250 x 20.2mm column. The column was eluted at 25 ml/min under the gradient

conditions of 30-60% acetonitrile in 5mM ammonium phosphate buffer over 25 min followed by holding at 60% acetonitrile in 5mM ammonium phosphate buffer for another 10 min. The desired fractions were collected, the acetonitrile was removed by evaporation and the residue was applied to a Bondesil 40µ C8 resin column. The column was washed with water and the product was eluted with methanol. Evaporation of the methanol gave compound 253 as a pale yellow solid.

EXAMPLE 1g - PREPARATION OF COMPOUND 76 and 177

Daptomycin (1ml, 0.1 M in dry dimethylformamide) was treated successively with 4-phenylbenzaldehyde (0.2 ml, 0.5M in dry dimethylformamide) and sodium triacetoxyborohydride (0.4 ml, 1.5M in dry dimethylformamide). The reaction mixture was capped and shaken briefly to mix the solution. After 24h the mixture was treated with water (0.2 ml) and loaded on an IBSIL-C8 5μ 250 x 20.2mm column. The column was eluted at 25 ml/min under the gradient conditions of 30-60% acetonitrile in 5mM ammonium phosphate buffer over 25 min followed by holding at 60% acetonitrile in 5mM ammonium phosphate buffer for another 10 min. The desired fractions were collected, the acetonitrile was removed by evaporation and the residue was applied to a Bondesil 40μ C8 resin column. The column was washed with water and the product was eluted with methanol. Evaporation of the methanol gave compound 76 as a pale yellow solid. Compound 177 was obtained by deacylation of compound 76 according to Example 7.

EXAMPLE Ih - PREPARATION OF COMPOUND 209

4-Hydroxy-3-nitrobenzaldehyde (0.4 ml, 0.2M in acetone) was successively treated with potassium hydroxide (0.1 ml, 1M in water) and 4-fluorobenzylbromide (0.4ml, 0.2M in acetone). After 24h the mixture was dried in vacuo to give 4-(4-fluorobenzyloxy)-3-nitro-benzaldehyde which was carried forward without further purification.

Daptomycin (1ml, 0.1 M in dry dimethylformamide) was treated successively with, 4-(4-fluorobenzyloxy)-3-nitro-benzaldehyde previously prepared above diluted in dry dimethylformamide (0.2 ml), and sodium triacetoxyborohydride

(0.4 ml, 1.5M in dry dimethylformamide). The mixture was capped and shaken briefly. After 24h the mixture was treated with water (0.2 ml) and loaded onto an IBSIL-C8 5µ 250 x 20.2mm column. The column was eluted at 25 ml/min under the gradient conditions of 30-60% acetonitrile in 5mM ammonium phosphate buffer over 25 min followed by holding at 60% acetonitrile in 5mM ammonium phosphate buffer for another 10 min. The desired fractions were collected, the acetonitrile was removed by evaporation and the residue was applied to a Bondesil 40µ C8 resin column. The column was washed with water and the product was eluted with methanol. Evaporation of the methanol gave compound 209 as a pale yellow solid.

EXAMPLE 2 - PREPARATION OF COMPOUNDS 10, 11-17, 19-20, 22-27 and 190

Daptomycin (972 mg) was dissolved in dry dimethylformamide (20 ml), and isatoic anhydride (979 mg) was added. The mixture was stirred at ambient temperature for 10 days, then quenched by the addition of water (20ml). The mixture was loaded onto a Bondesil 40μ C8 resin column (25g), which had been previously washed with methanol (50 ml) and water (100ml). The column was then eluted with water (200ml), 15% methanol/water (1200ml), 20% methanol/water (200ml), 30% methanol/water (200ml) and 40% methanol/water (200ml). The product bearing fractions were combined and freeze dried to give compound 10 as a white solid (870 mg).

In an analogous manner, compounds 11-17, 19-20, 22-27 and 190 can be prepared as detailed in the above example by appropriate substitutions of reagents obvious to those skilled in the art following the teachings of the disclosure.

EXAMPLE 3 - PREPARATION OF COMPOUNDS 44, 45, 41-43, 46-48, 55-58, 60-75, 178-180, 193 and 245

Daptomycin (500 mg) and Boc-tryptophan-p-nitrophenyl ester (157.5 mg) were stirred at room temperature in dry dimethylformamide (30 ml) for 3 days. Water (30 ml) was added and the mixture was purified on a Bondesil 40µ C8 resin column (25 g). The column was eluted with 20% acetonitrile in water (200 ml), 40% acetonitrile in water (200 ml) and finally with methanol. Evaporation of the solvent

from the product-containing fractions gave compound 44 as a pale yellow solid (450 mg).

Compound 44 (200 mg) was cooled to 0°C and a 0°C solution of 5% thioanisole in trifluoroacetic acid (10 ml) was added. After 3 hours at 0°C the mixture was evaporated to dryness and the residue was purified by preparative HPLC on an IBSIL-C8 5µ 250x20.2mm column. The column was eluted at 20 ml/min with 38% acetonitrile in 5mM ammonium phosphate buffer. The product containing fractions were freeze-dried. The freeze-dried residue was dissolved in water (5 ml) and applied to a Bondesil 40µ C8 resin column. The column was washed with water and eluted with methanol. Evaporation of the methanol gave compound 45 as a pale yellow solid.

In an analogous manner, compounds 41-43, 46-48, 55-58, 60-75, 178-180, 193 and 245 can be prepared as detailed in the above example by appropriate substitutions of reagents obvious to those skilled in the art following the teachings of the disclosure.

EXAMPLE 3a - PREPARATION OF COMPOUNDS 54, 49 and 51

Daptomycin (400 mg) and N, N-bis(tert-butoxycarbonyl)-L-lysine-4-nitrophenyl ester (173 mg) were stirred in dry dimethylformamide (5 ml) at room temperature for two days. The mixture was loaded onto an IBSIL-C8 5 μ 250x20.2 mm column and was eluted at 20 ml/min with 37% acetonitrile in 5 mM ammonium phosphate buffer. Fractions containing the desired compound were collected and freeze-dried. The freeze-dried residue was dissolved in water (5 ml) and applied to a Bondesil 40 μ C8 resin column, washed with water and eluted with methanol. Evaporation of the methanol gave the Boc protected intermediate as a pale yellow solid (370 mg).

Boc protected intermediate (200 mg) was stirred in trifluoroacetic acid (5 ml) and anisole (0.25 ml) at room temperature for 2 hours. Removal of the solvents under reduced pressure gave a residue which was loaded on an IBSIL-C8 5 μ 250x20.2 mm column and eluted at 20 ml/min with 37% acetonitrile in 5 mM ammonium phosphate buffer. Fractions containing the desired compound were

collected and freeze-dried. The freeze-dried residue was dissolved in water (5 ml) and applied to a Bondesil 40 μ C8 resin column, washed with water and eluted with methanol. Evaporation of the methanol gave compound 54 as a pale yellow solid (100 mg).

In an analogous manner, compounds 49 and 51 can be prepared as detailed in the above example by appropriate substitutions of reagents obvious to those skilled in the art following the teachings of the disclosure.

EXAMPLE 3b - PREPARATION OF COMPOUNDS 32, 18, 21, 28-31, 33-35, 39, 182 and 189

Daptomycin (162 mg) and 2-methylthiobenzoic acid pentafluorophenol ester (37 mg) were stirred at room temperature in dry dimethylformamide (10 ml) for 5 days. The dimethylformamide was evaporated under reduced pressure and the residue was purified by preparative HPLC on an IBSIL-C8 5μ 250x20.2mm column. The column was eluted at 20 ml/min with 36% acetonitrile in 5mM ammonium phosphate buffer. Fractions collected at 7.3 minutes were freeze-dried. The freeze-dried residue was dissolved in water (5 ml) and applied to a Bondesil 40μ C8 resin column. The column was washed with water and eluted with methanol. Evaporation of the methanol gave compound 32 as a pale yellow solid (47 mg).

In an analogous manner, compounds 18, 21, 28-31, 33-35, 39, 182 and 189 can be prepared as detailed in the above example by appropriate substitutions of reagents by one having ordinary skill in the art following the teachings of the disclosure.

EXAMPLE 4 - PREPARATION OF COMPOUNDS 5, 4, 6-8 and 9

Daptomycin (16 mg) was dissolved in dry dimethylformamide (0.5 ml) and methyl isothiocyanate (37 mg) was added. The mixture was stirred at ambient temperature for 24 hours, then quenched by the addition of 5% ammonium phosphate buffer (1ml). The mixture was purified by preparative HPLC on an IBSIL-C8 5μ 250x20.2mm column. The column was eluted at 20 ml/min with 36% acetonitrile in 5 mM ammonium phosphate buffer. The product bearing fractions were combined and

freeze dried. The freeze-dried residue was dissolved in water (1.5 ml) and applied to a Bondesil 40μ C8 resin column. The column was washed with water and eluted with methanol. Evaporation of the methanol gave compound 5 as a pale yellow solid (5.2 mg).

In an analogous manner, compounds 4, 6-8 and 9 can be prepared as detailed in the above example by appropriate substitutions of reagents obvious to those having ordinary skill in the art.

EXAMPLE 5 - PREPARATION OF COMPOUND 3

Daptomycin (16 mg) and N-benzotriazole phenylsulfonamide (2.6 mg) were stirred at room temperature in dry pyridine for 6 days. The solvent was evaporated and the residue was purified by preparative HPLC using an IBSIL-C8 5μ 250x20.2mm column. The column was eluted at 20 ml/min with 36% acetonitrile in 5mM ammonium phosphate buffer and product containing fractions were freezedried. The freeze dried residue was dissolved in water (5 ml) and applied to a Bondesil 40μ C8 resin column. The column was washed with water and eluted with methanol. Evaporation of the methanol gave compound 3 as a pale yellow solid (4 mg).

EXAMPLE 6 - PREPARATION OF COMPOUNDS 1, 2, 221, 259 and 270

Daptomycin (32 mg) was dissolved in dry dimethylformamide (20 ml), and N,N'-bis-Boc-1-guanidinylpyrazole (31 mg) was added. The mixture was stirred at ambient temperature for 5 days, then quenched by the addition of water (3ml). The resultant mixture was loaded onto a Bondesil 40μ C8 resin (900 mg) that had been previously washed with methanol and water. The column was eluted with water (30ml) followed by methanol. The product-bearing fractions were combined and evaporated to give compound 1 as a white solid.

Compound 1 (30 mg) was dissolved in trifluoroacetic acid/dichloromethane/tri-isopropylsilane/ethane dithiol (11/8/0.5/0.5, 3ml) and stirred at ambient temperature for 90 minutes. The mixture was evaporated to dryness and the residue was precipitated by the addition of diethyl ether (10 ml). The residue was

purified by preparative HPLC on an IBSIL-C8 5µ 250x20.2mm column. The column was eluted at 20 ml/min with 38% acetonitrile in 5 mM ammonium phosphate buffer. The product-bearing fractions were combined and freeze dried. The freeze-dried residue was dissolved in water (1.5 ml) and applied to a Bondesil 40µ C8 resin column. The column was washed with water and eluted with methanol. Evaporation of the methanol gave compound 2 as a white solid (6.4 mg).

In an analogous manner, compounds 221, 259 and 270 can be prepared as detailed in the above example by appropriate substitutions of reagents obvious to those having ordinary skill in the art following the teachings of the disclosure.

EXAMPLE 7 - PREPARATION OF COMPOUNDS 255, 260, 254, 256-257, 261, 263, 292-294 and 313-314

Daptomycin (10g) was dissolved in dry dimethylformamide (100ml). N,N'-bis-Boc-guanidinylpyrazole (2.3g) in dry dimethylformamide (5ml) was added. The mixture was stirred under nitrogen at room temperature overnight. The mixture was purified on a Bondesil 40µ C8 resin column. The product containing fractions were freeze-dried to give compound 1 (7.4g) as pale yellow fluffy solid.

Compound 1 (2.6g) was added to a preparation of deacylase enzyme produced from recombinant Streptomyces lividans, which expresses the Actinoplanes utahensis deacylase enzyme in ethylene glycol (1.2 ml) and water (25 ml). The pH of the solution was adjusted to 9 with 1.0M sodium hydroxide solution and stirred at room temperature. After 24 hours the mixture was purified on a Bondesil 40µ C8 resin column by eluting with 10% acetonitrile/water, then 40% acetonitrile/water. The product-containing fractions were freeze dried to give deacylated bis-Bocguanidinylated daptomycin (0.69 g) as a pale yellow solid.

Undecanoyl pentafluorophenol ester (40.3 mg) was added to deacylated bis-Boc-guanidinylated daptomycin (171.5 mg) in dry dimethylformamide (2 ml). The mixture was stirred overnight at room temperature before being concentrated to give compound 255 (105 mg) as a yellow solid.

Compound 255 was dissolved in trifluoroacetic acid (5.5 ml), dichloromethane (4 ml), ethane dithiol (0.25 ml) and triisopropylsilane (0.25ml). The

mixture was stirred for 4 hours at room temperature before being concentrated and purified by preparative HPLC on an IB-SIL 5µ 250x20.2mm column. The column was eluted at 25 ml/min with acetonitrile and ammonium phosphate buffer 30%-60% gradient for 40 min. The desired fractions were collected at 21 minutes and freeze dried. The freeze-dried residue was dissolved in water and applied to a Bondesil C8 resin column. The column was washed with water and eluted with methanol. Evaporation of the methanol gave compound 260 (27.8 mg) as a pale yellow solid.

In an analogous manner, compounds 254, 256-257, 261, 263, 292-294 and 313-314 can be prepared as detailed in the above example by appropriate substitutions of reagents obvious to those having ordinary skill in the art following the disclosure of the invention.

EXAMPLE 7a - PREPARATION OF COMPOUNDS 258 and 262

Tetradecanoyl pentafluorophenol ester (35.5 mg) and deacylated bis-Boc-guanidinylated daptomycin (102.5 mg) in dry dimethylformamide (2 ml). The mixture was stirred overnight at room temperature before being concentrated to give compound 258 (38.8 mg) as a yellow solid.

Compound 258 (38.8 mg) was dissolved in trifluoroacetic acid (5.5 ml), dichloromethane (4 ml), ethane dithiol (0.25 ml) and triisopropylsilane (0.25 ml). The mixture was stirred for 4 hours at room temperature before being concentrated and purified by preparative HPLC on an IB-SIL 5µ 250x20.2mm column. The column was eluted at 25 ml/min with acetonitrile and ammonium phosphate buffer 30%-60% gradient for 40 min. The desired fractions were collected at 21minutes and freeze dried. The freeze-dried residue was dissolved in water and applied to a Bondesil C8 resin column. The column was washed with water and eluted with methanol. Evaporation of the methanol gave compound 262 (2.1 mg) as a pale yellow solid.

EXAMPLE 8 - PREPARATION OF COMPOUND 37, 36 and 192

Daptomycin (162 mg) was stirred in 0.1 M hydrochloric acid (5 ml) at 0°C for 10 minutes before sodium nitrite (8 mg) in water (0.2 ml) was added

dropwise. Sulfamic acid (11 mg) was added after 15 minutes, followed by sodium azide (8 mg) 10 minutes later. The mixture was maintained at 0°C for 4 hours and then neutralized with a saturated sodium bicarbonate solution and purified by preparative HPLC. An IBSIL-C8 5µ 250x20.2mm column was loaded with the mixture and eluted at 20 ml/min with 37% acetonitrile in 5mM ammonium phosphate buffer. Fractions were collected at 6.9 minutes and freeze dried. The freeze-dried residue was dissolved in water (5 ml) and applied to a Bondesil 40µ C8 resin column. The column was washed with water and eluted with methanol. Evaporation of the methanol gave the azido daptomycin as a pale yellow solid (60 mg).

The azido daptomycin (69 mg) was dissolved in dry dimethylformamide (4 ml) and iminobiotin-N-hydroxysuccinimide ester (53 mg) was added. The mixture was covered to exclude light and stirred at ambient temperature for 3 days. The mixture was quenched by the addition of water (20ml). The resultant mixture was loaded onto a Bondesil 40µ C8 resin (25g) column, which had been previously washed with methanol and water, and the column was eluted with water. The product-bearing fractions were combined and freeze dried to give Compound 37 as a white solid (49 mg).

In an analogous manner, compounds 36 and 192 can be prepared as detailed in the above example by appropriate substitutions of reagents obvious to those having ordinary skill in the art by following the disclosure of the invention.

EXAMPLE 8a - PREPARATION OF COMPOUND 200

Daptomycin (1.62 g) in 50% wt aqueous solution of hypophosphorus acid (10 ml) was stirred at 0°C for 30 minutes before adding dropwise a solution of sodium nitrite (76 mg) in water (0.5 ml). The mixture was allowed to come to room temperature and stirred for 24 hours. The mixture was purified by preparative HPLC by loading the mixture on an IBSIL-C8 5μ 250x20.2mm column and eluting the column at 20 ml/min with 32% acetonitrile in 5mM ammonium phosphate buffer. The desired fractions were collected at 30 minutes and freeze dried. The freeze-dried residue was dissolved in water (5 ml) and applied to a Bondesil 40μ C8 resin column.

The column was washed with water and eluted with methanol. Evaporation of the methanol gave desamino daptomycin as a pale yellow solid (200 mg).

To desamino daptomycin (80 mg) in dry dimethylformamide (2 ml) was added N-t-butoxycarbonyl-L-tryptophan-p-nitrophenyl ester (32 mg). The mixture was stirred at room temperature for 24 hours before being purified by preparative HPLC. The mixture was loaded on an IBSIL-C8 5µ 250x20.2mm column and eluted at 20 ml/min with 40 % acetonitrile in 5mM ammonium phosphate buffer. The desired fractions were collected at 19 minutes and freeze-dried. The freeze-dried residue was dissolved in water (2 ml) and applied to a plug of Bondesil 40µ C8 resin (500 mg). The Bondesil resin was washed with water (10 ml) and then the product was eluted with methanol (10 ml). Evaporation of the methanol gave Boc protected compound 200 as a pale yellow solid (20 mg).

To Boc protected compound 200 (20 mg) in 60% trifluoroacetic acid in dichloromethane (0.5 ml) was added anisole (10 μ L). The mixture was stirred at room temperature for 6 hours before being evaporated to dryness. Preparative HPLC purification of the residue was done on an IBSIL-C8 5 μ 250x20.2mm column and eluted at 20 ml/min with 38 % acetonitrile in 5mM ammonium phosphate buffer. The desired fractions were collected at 15 minutes and freeze-dried. The freeze-dried residue was dissolved in water (2 ml) and applied to a plug of Bondesil 40 μ C8 resin (500 mg). The Bondesil resin was washed with water (10 ml) and the product was eluted with methanol (10 ml). Evaporation of the methanol gave compound 200 as a pale yellow solid (4 mg).

EXAMPLE 9 - PREPARATION OF COMPOUNDS 181, 86, 101-102, 206-207, 213-220, 246-251, 264 and 269

Daptomycin (250 mg) and N-tBoc-L-tryptophan-p-nitrophenyl ester (144 mg) were stirred in dry dimethylformamide (3 ml) at room temperature for two days. The mixture was loaded on an IBSIL-C8 5 μ 250x20.2 mm column and was eluted at 20 ml/min with 37% acetonitrile in 5 mM ammonium phosphate buffer. Fractions containing the desired compound were collected and freeze-dried. The freeze-dried residue was dissolved in water (5 ml) and applied to a Bondesil 40 μ C8

resin column, washed with water and eluted with methanol. Evaporation of the methanol gave N-Boc tryptophan daptomycin as a pale yellow solid (130 mg).

A preparation of deacylase enzyme was produced from recombinant *Streptomyces lividans*, which expresses the *Actinoplanes utahensis* deacylase enzyme. The enzyme in ethylene glycol (400 µl) was added to the solution of N-Boc tryptophan daptomycin (100 mg) in HPLC grade water (20 ml). The solution was adjusted to pH 8.5 with sodium hydroxide (1 M). The mixture was stirred for 24 hours. The mixture was loaded on a C8 resin plug column, washed with water and eluted with methanol. Evaporation of the methanol gave a residue which was applied to an IBSIL-C8 5 µ 250x20.2 mm column and was eluted at 20 ml/min with 37% acetonitrile in 5 mM ammonium phosphate buffer. Fractions containing the desired compound were collected and freeze-dried. The freeze-dried residue was dissolved in water (5 ml) and applied to a Bondesil 40 µ C8 resin column, washed with water and eluted with methanol. Evaporation of the methanol gave deacylated N-Boc tryptophan daptomycin as a pale yellow solid (42 mg).

Deacylated N-Boc tryptophan daptomycin (20 mg) was stirred in dry dimethylformamide (2 ml) at room temperature. Undecyl isocyanate (2.25 mg) was added to the solution. After stirring at ambient temperature for 24 hours, the mixture was diluted with water (10 ml) and applied to a Bondesil 40 μ C8 resin column, washed with water and eluted with methanol. Evaporation of the methanol gave the undecyl urea of N-Boc tryptophan daptomycin as a pale yellow solid (21 mg).

N-Boc tryptophan daptomycin undecyl urea (21 mg) was stirred in trifluoroacetic acid (2 ml) and anisole (0.1 ml) at room temperature for 2 hours. Removal of the solvents under reduced pressure gave a residue which was loaded on an IBSIL-C8 5 μ 250x20.2 mm column and eluted at 20 ml/min with 37% acetonitrile in 5 mM ammonium phosphate buffer. Fractions containing the desired compound were collected and freeze-dried. The freeze-dried residue was dissolved in water (5 ml) and applied to a Bondesil 40 μ C8 resin column, washed with water and eluted with methanol. Evaporation of the methanol gave compound 181 as a pale yellow solid (0.8 mg).

In an analogous manner, compounds 86, 101-102, 206-207, 213-220, 246-251, 264 and 269 can be prepared as detailed in the above example by appropriate substitutions of reagents obvious to those having ordinary skill in the art following the disclosure of the invention.

EXAMPLE 9a - PREPARATION OF COMPOUND 205

Deacylated N-Boc tryptophan daptomycin (50 mg) and nonaldehyde (4.1 mg) were stirred in dry dimethylformamide (2 ml) at room temperature. Sodium triacetoxy borohydride (3.6 mg) was added to the solution. The mixture was stirred for 24 hours, then loaded on an IBSIL-C8 5 μ 250x20.2 mm column and eluted at 20 ml/min with 37% acetonitrile in 5 mM ammonium phosphate buffer. Fractions containing the desired compound were collected and freeze-dried. The freeze-dried residue was dissolved in water (5 ml) and applied to a Bondesil 40 μ C8 resin column, washed with water and eluted with methanol. Evaporation of the methanol gave nonyl amino N-Boc tryptophan daptomycin as a pale yellow solid (14 mg).

Nonyl amino N-Boc tryptophan daptomycin (14 mg) was stirred in trifluoroacetic acid (2 ml) and anisole (0.1 ml) at room temperature for 2 hours. Removal of the solvents under reduced pressure gave a residue which was loaded on an IBSIL-C8 5 μ 250x20.2 mm column and was eluted at 20 ml/min with 37% acetonitrile in 5 mM ammonium phosphate buffer. Fractions containing the desired compound were collected and freeze-dried. The freeze-dried residue was dissolved in water (5 ml) and applied to a Bondesil 40 μ C8 resin column, washed with water and eluted with methanol. Evaporation of the methanol gave compound 7 as a pale yellow solid (5 mg).

EXAMPLE 10 - PREPARATION OF COMPOUNDS 356, 315-322, 332-337, 345-349 and 355

Daptomycin (5.0 g) was dissolved in water (25 ml) and adjusted to pH 9 with 5M sodium hydroxide. Di-tert-butyldicarbonate (1.5 g) was added and the mixture was adjusted to maintain pH 9 with 5 M sodium hydroxide until the reaction was complete (4 hours). The pH was adjusted to 7 and the mixture was loaded onto a

Bondesil 40µ C8 resin column. The column was washed with water and the product was eluted from the column with methanol. Evaporation of the methanol gave Bocprotected daptomycin (5.08 g) as a yellow powder.

A preparation of deacylase enzyme was produced from recombinant Streptomyces lividans, which expresses the Actinoplanes utahensis deacylase enzyme. The enzyme in ethylene glycol (400 µl) was added to Boc-protected daptomycin (1 g) in water (100 ml) at pH 7-8. After incubation for 72 hours, the mixture was loaded on a Bondesil 40µ C8 resin column. The column was washed with water and the product was eluted from the column with 10% acetonitrile in water. The solvent was removed by evaporation to give deacylated Boc-protected daptomycin (440 mg) as a yellow powder.

Daptomycin undecyl urea synthesized from deacylated Boc protected daptomycin above using undecyl isocyanate instead of undecanoyl pentafluorophenol ester according to example 7 (100mg) and 5-methoxyindole-3-carboxaldehyde (11mg) in dry dimethylformamide (0.6ml) was added sodium triacetoxyborohydride (76mg). The mixture was stirred at room temperature for 24 hours before purification by preparative HPLC. The mixture was loaded on an IBSIL-C8 5µ 250x20.2mm column and eluted at 25 ml/min with 30-60% acetonitrile in 5mM ammonium phosphate gradient over 30 minutes. The desired fractions were collected at 21 minutes and freeze-dried. The freeze-dried residue was dissolved in water (2ml) and applied to a plug of Bondesil 40µ C8 resin (500mg). The Bondesil resin was washed with water (10ml) and then the product was eluted with methanol (10ml). Evaporation of the methanol gave compound 114 as a pale yellow solid (10mg).

In an analogous manner, compounds 315-322, 332-337, 345-349 and 355 can be prepared as detailed in the above example by appropriate substitutions of reagents obvious to those skilled in the art.

EXAMPLE 10a - PREPARATION OF COMPOUNDS 307, 310, 295-306, 308-309, 311-312, 338-344 and 350-352

Daptomycin undecanoyl amide synthesized from deacylated Boc protected daptomycin by using undecanoyl pentafluorophenol ester according to

examples 10 and 7 (60 mg) was stirred in dry dimethylformamide (2 ml) at room temperature. N-tBoc-L-tryptophan-p-nitrophenyl ester (31 mg) was added to the solution. The mixture was stirred for 24 hours. The mixture was loaded onto an IBSIL-C8 5 μ 250x20.2 mm column and was eluted at 20 ml/min with 37% acetonitrile in 5 mM ammonium phosphate buffer. Fractions containing the desired compound were collected and freeze-dried. The freeze-dried residue was dissolved in water (5 ml) and applied to a Bondesil 40 μ C8 resin column, washed with water and eluted with methanol. Evaporation of the methanol gave compound 307 as a pale yellow solid (25 mg).

Compound 307 (20 mg) was stirred in trifluoroacetic acid (2 ml) and anisole (0.1 ml) at room temperature for 2 hours. Removal of the solvents under reduced pressure gave a residue which was loaded on an IBSIL-C8 5 μ 250x20.2 mm column and was eluted at 20 ml/min with 37% acetonitrile in 5 mM ammonium phosphate buffer. Fractions containing the desired compound were collected and freeze-dried. The freeze-dried residue was dissolved in water (5 ml) and applied to a Bondesil 40 μ C8 resin column, washed with water and eluted with methanol. Evaporation of the methanol gave compound 310 as a pale yellow solid (4 mg).

In an analogous manner, compounds 295-306, 308-309, 311-312, 338-344 and 350-352 can be prepared as detailed in the above example by appropriate substitutions of reagents obvious to those skilled in the art.

EXAMPLE 11

Compounds according to Formula I were tested for antimicrobial activity against a panel of organisms according to standard procedures described by the National Committee for Clinical Laboratory Standards (NCCLS document M7-A5, Vol. 20, No. 2, 2000) except that all testing was performed at 37°C. Compounds were dissolved in 100% dimethyl sulfoxide and were diluted to the final reaction concentration (0.1 μ g/mL-100 μ g/mL) in microbial growth media. In all cases the final concentration of dimethyl sulfoxide incubated with cells is less than or equal to 1%. For minimum inhibitory concentration (MIC) calculations, 2-fold dilutions of compounds were added to wells of a microtiter plate containing $5x10^4$

bacteria cells in a final volume of $100~\mu L$ of media (Mueller-Hinton Broth supplemented with 50~mg/L Ca^{2+}). The optical densities (OD) of the bacterial cells, which measures bacterial cell growth and proliferation, were measured using a commercial plate reader. The MIC value is defined as the lowest compound concentration inhibiting growth of the test organism. The MIC (in $\mu g/ml$) values of representative compounds of the present invention are listed in Table III.

EXAMPLE 12

The mouse protection test is an industry standard for measuring the efficacy of a test compound in vivo [for examples of this model see J. J. Clement, et al., Antimicrobial Agents and Chemotherapy, 38 (5), 1071-1078, (1994)]. As exemplified below, this test is used to demonstrate the in vivo efficacy of the compounds of the present invention against bacteria.

The *in vivo* antibacterial activity was established by infecting female CD-1 mice (Charles River Lab, MA) weighing 19–23 g intraperitoneally with from Methicillin Resistant *S. aureus* (MRSA) inoculum. The inoculum was prepared from Methicillin Resistant *S. aureus* (ATCC 43300). The MRSA inoculum was cultured in Mueller-Hinton (MH) broth at 37° C for 18 hours. The optical density at 600 nm (OD₆₀₀) was determined for a 1:10 dilution of the overnight culture. Bacteria (8 x 10⁸ cfu) was added to 20 ml of phosphate buffered saline (Sigma P-0261) containing 5 % hog gastric mucin (Sigma M-2378). All animals were injected with 0.5 ml of the inoculum, equivalent to 2 x 10⁷ cfu/mouse, which is the dose causing ~100% death of the animals without treatment.

The test compound was dissolved in 10.0 ml of 50mM phosphate buffer to give a solution of 1 mg/ml (pH = 7.0). This solution was serially diluted with vehicle by 4-fold (1.5 ml to 6.0 ml) to give 0.25, 0.063 and 0.016 mg/ml solutions. All the solutions were filtered with 0.2 m Nalgene syringe filter. Immediately after the bacterial inoculation, group 1 animals were subcutaneously (sc) injected with buffer (no test compound) and groups 2 to 5 were given test compound sc at 10.0, 2.5, 0.63, and 0.16 mg/kg, respectively. Group 6 animals

received test compound sc at 10 mg/kg (or the highest therapeutic dose of a given compound) only for monitoring acute toxicity. These injections were repeated once at 4 hours after the inoculation for the respective groups. The injection volume at each time was 10 ml per kilogram of body weight. The results of the *in vivo* efficacy test are summarized in Table II, which provides a representative example of the results obtained for Compound 70. The 50% effective dose (ED₅₀) is calculated on the basis of the number of mice surviving 7 days after inoculation. The ED₅₀ was determined for other compounds of this invention in a similar manner. The ED₅₀ in mg/kg of other representative compounds of the present invention are listed in Table III.

Table II

Group	# of mice	Inoculated with	Treatment	Survival (7 days)
1	5	MRSA #43300 2x10 ⁷ cfu/mouse	Phosphate buffer 10 ml/kg, s.c. x2	0/5
2	5	<i>MRSA #43300</i> 2x10 ⁷ cfu/mouse	Compound 70 10 mg/kg, s.c. x2	5/5
3	5	MRSA #43300 2x10 ⁷ cfu/mouse	Compound 70 2.5 mg/kg, s.c. x2	3/5
4	5	MRSA #43300 2x10 ⁷ cfu/mouse	Compound 70 0.63 mg/kg, s.c. x2	1/5
5	5	MRSA #43300 2x10 ⁷ cfu/mouse	Compound 70 0.16 mg/kg, s.c. x2	0/5
6	5	No	Compound 70 10 mg/kg, s.c. x2	5/5

The ED₅₀ of compound 70 is calculated to be 1.51 mg/kg

Table III

	MIC	MIC	ED ₅₀
Compound #	(µg/ml)	(μg/ml)	mg/kg
_	S. aureus	E. faecalis	S. aureus
1	++	+	++
2	+++	+	+++
3	++	+	
4	+	+	
5	++	++	
6	++	++	
7	++	++	
8	++	++	
9	+++	++	
10	+++	+	++
11	++	+	
12	+++	++	
13	+++	++	
14	++	++	
15	++	1-1	
16	+++	++.	
17	++	++	
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Wherein "+++" indicates that the compound has an MIC (μ g/ml) of 1 μ g/ml or less or an ED₅₀ of 1 mg/kg or less;

"++" indicates that the compound has an MIC ($\mu g/ml$) or ED₅₀ of greater than 1 $\mu g/ml$ or 1 mg/kg, respectively but less than or equal to 10 $\mu g/ml$ or ED₅₀ of 10 mg/kg, respectively; and

"+" indicates that the compound has an MIC (μ g/ml) of greater than 10 μ g/ml or an ED₅₀ of greater than 10 mg/kg.

All publications and patent applications cited in this specification are herein incorporated by reference as if each individual publication or patent application were specifically and individually indicated to be incorporated by reference. Although the foregoing invention has been described in some detail by way of illustration and example for purposes of clarity of understanding, it will be readily apparent to those of ordinary skill in the art in light of the teachings of this invention that certain changes and modifications may be made thereto without departing from the spirit or scope of the appended claims.

CLAIMS

We claim:

1. A compound having the formula (I):

and salts thereof;

wherein R is:

wherein X and X" are independently selected from C=O, C=S, C=NH, C=NR X , S=O or SO₂,

wherein n is 0 or 1;

wherein R^X is selected from alkyl, alkenyl, alkynyl, aryl, heteroaryl, cycloalkyl, heterocyclyl, hydroxyl, alkoxy, carboxy or carboalkoxy,

wherein B is X"RY, H, alkyl, alkenyl, alkynyl, aryl, heteroaryl, cycloalkyl or heterocyclyl;

wherein R^Y is selected from hydrido, alkyl, alkenyl, alkynyl, aryl, heteroaryl, cycloalkyl, heterocyclyl or hydroxyl,

wherein A is H, NH₂, NHR^A, NR^AR^B, alkyl, alkenyl, alkynyl, alkoxy, aryloxy, aryl, heteroaryl, cycloalkyl or heterocyclyl;

wherein R^A and R^B are independently selected from alkyl, alkenyl, alkynyl, aryl, heteroaryl, cycloalkyl, heterocyclyl or carboalkoxy;

wherein when n is 0, then A is additionally selected from:

wherein each of R⁵⁰-R⁵³ is independently selected from C₁-C₁₅ alkyl, alternatively, wherein B and A together form a 5-7 membered heterocyclic or heteroaryl ring;

wherein R¹ is

wherein X' and X''' are independently selected from C=0, C=S, C=NH, C=NR $^{X'}$, S=O or SO₂,

wherein m is 0 or 1;

wherein R^{X'} is selected from alkyl, alkenyl, alkynyl, aryl, heteroaryl, cycloalkyl, heterocyclyl, hydroxyl, alkoxy, carboxy or carboalkoxy,

wherein B' is $X'''R^{Y'}$, H, alkyl, alkenyl, alkynyl, aryl, heteroaryl, cycloalkyl or heterocyclyl,

wherein R^{Y} is selected from hydrido, alkyl, alkenyl, alkynyl, aryl, heteroaryl, cycloalkyl, heterocyclyl or hydroxyl;

wherein A' is H, NH₂, NHR^{A'}, NR^{A'}R^{B'}, heteroaryl, cycloalkyl or heterocyclyl,

wherein R^{A'} and R^{B'} are independently selected from alkyl, alkenyl, alkynyl, aryl, heteroaryl, cycloalkyl, heterocyclyl or carboalkoxy,

wherein when m is 0, then A' is additionally selected from:

$$- \left\{ \begin{array}{c} O \\ P \\ OR^{50} \end{array} \right. \quad \left\{ \begin{array}{c} O \\ P \\ R^{53} \end{array} \right. \quad \text{and} \quad \left\{ \begin{array}{c} O \\ P \\ R^{53} \end{array} \right. \quad \text{and} \quad \left\{ \begin{array}{c} O \\ P \\ R^{53} \end{array} \right.$$

wherein each of R^{50} - R^{53} is independently selected from C_1 - C_{15} alkyl, provided that when B' is H and X' is C=O, then A' is other than

- (a) a pyridinyl ring substituted with one substitutent NHC(O)R^D or
- (b) a C₅-C₆ saturated cycloalkyl ring substituted with one substitutent

 $NHC(O)R^{D}$;

wherein R^D is C_1 - C_{17} unsubstituted alkyl or C_2 - C_{17} unsubstituted

alkenyl; and

when B' is H and m=0, then A' is not H; wherein R² is

wherein K and K' together form a C_3 - C_7 cycloalkyl or heterocyclyl ring or a C_5 - C_{10} aryl or heteroaryl ring,

wherein J is selected from the group consisting of hydrido, amino, NHR^J, NR^JR^K, alkyl, alkenyl, alkynyl, alkoxy, aryloxy, aryl, heteroaryl, cycloalkyl, heterocyclyl, alkylamino, hydroxyl, thio, alkylthio, alkenylthio, sulfinyl, sulfonyl, azido, cyano, halo,

$$- \begin{cases} S \\ NR^{24}R^{25} \end{cases} \text{ and } - \begin{cases} S \\ OR^{26} \end{cases}$$

wherein each of R²⁴, R²⁵, and R²⁶ is independently selected from the group consisting of alkyl, cycloalkyl, heterocyclyl, aryl and heteroaryl; or R²⁴ and R²⁵ together form a 5-8 membered heterocyclyl ring;

wherein R^I and R^K are independently selected from alkyl, alkenyl, alkynyl, aryl, heteroaryl, cycloalkyl or heterocyclyl, or

alternatively, wherein J, together with R^{17} , forms a 5-8 membered heterocyclyl or cycloalkyl ring; or

alternatively, wherein J, together with both R¹⁷ and R¹⁸, forms a 5-8 membered aryl, cycloalkyl, heterocyclyl or heteroaryl ring, and

wherein each of R¹⁷ and R¹⁸ is independently selected from the group consisting of hydrido, halo, hydroxyl, alkoxy, amino, thio, sulfinyl, sulfonyl and

wherein R^{17} and R^{18} taken together can form a group consisting of ketal, thioketal,

$$= \begin{cases} = 0 & , = \\ = S & , = \\ = NOR^{22} \text{ and } = \end{cases} = NNR^{22}R^{23}$$

wherein each of R^{22} and R^{23} is independently selected from the group consisting of hydrido and alkyl.

2. A compound having the formula (I):

and salts thereof;

wherein R is:

wherein X and X" are independently selected from C=O, C=S, C=NH, C=NR^X, S=O or SO₂;

wherein n is 0 or 1;

wherein R^X is selected from alkyl, alkenyl, alkynyl, aryl, heteroaryl, cycloalkyl, heterocyclyl, hydroxyl, alkoxy, carboxy or carboalkoxy;

wherein B is X"R", H, alkyl, alkenyl, alkynyl, aryl, heteroaryl, cycloalkyl or heterocyclyl;

wherein RY is selected from hydrido, alkyl, alkenyl, alkynyl, aryl, heteroaryl, cycloalkyl, heterocyclyl or hydroxyl;

wherein A is H, NH₂, NHR^A, NR^AR^B, alkyl, alkenyl, alkynyl, alkoxy, aryloxy, aryl, heteroaryl, cycloalkyl or heterocyclyl;

wherein R^A and R^B are independently selected from alkyl, alkenyl, alkynyl, aryl, heteroaryl, cycloalkyl, heterocyclyl or carboalkoxy;

wherein when n is 0, then A is additionally selected from:

wherein each of R⁵⁰-R⁵³ is independently selected from C₁-C₁₅ alkyl; alternatively, wherein B and A together form a 5-7 membered heterocyclic or heteroaryl ring;

wherein R¹ is

wherein X' and X''' are independently selected from C=0, C=S, C=NH, C=NR $^{X'}$, S=O or SO₂,

wherein m is 0 or 1:

wherein R^{X'} is selected from alkyl, alkenyl, alkynyl, aryl, heteroaryl, cycloalkyl, heterocyclyl, hydroxyl, alkoxy, carboxy or carboalkoxy,

wherein B' is X"'RY', H, alkyl, alkenyl, alkynyl, aryl, heteroaryl, cycloalkyl or heterocyclyl;

wherein R^{Y'} is selected from hydrido, alkyl, alkenyl, alkynyl, aryl, heteroaryl, cycloalkyl, heterocyclyl or hydroxyl;

wherein A' is aryl;

provided that when B' is H and X' is C=O, then A' is other than a phenyl ring substituted with substitutent NHC(O)R^D, wherein R^D is C_1 - C_{17} unsubstituted alkyl or C_2 - C_{17} unsubstituted alkenyl, wherein said phenyl ring may be further optionally substituted with 1-2 substituents independently selected from amino, nitro, C_1 - C_3 alkyl, hydroxyl, C_1 - C_3 alkoxy, halo, mercapto, C_1 - C_3 alkylthio, carbamyl or C_1 - C_3 alkyl carbamyl;

wherein R² is

wherein K and K' together form a C_3 - C_7 cycloalkyl or heterocyclyl ring or a C_5 - C_{10} aryl or heteroaryl ring;

wherein J is selected from the group consisting of hydrido, amino, NHR^J, NR^JR^K, alkyl, alkenyl, alkynyl, alkoxy, aryloxy, aryl, heteroaryl, cycloalkyl, heterocyclyl, alkylamino, hydroxyl, thio, alkylthio, alkenylthio, sulfinyl, sulfonyl, azido, cyano, halo,

$$- \begin{cases} S & \text{and} \\ - S & \text{OR}^{26} \end{cases}$$

wherein each of R^{24} , R^{25} , and R^{26} is independently selected from the group consisting of alkyl, cycloalkyl, heterocyclyl, aryl and heteroaryl, or R^{24} and R^{25} together form a 5-8 membered heterocyclyl ring,

wherein R^{J} and R^{K} are independently selected from alkyl, alkenyl, alkynyl, aryl, heteroaryl, cycloalkyl or heterocyclyl; or

alternatively, wherein J, together with R¹⁷, forms a 5-8 membered heterocyclyl or cycloalkyl ring, or

alternatively, wherein J, together with both R^{17} and R^{18} , forms a 5-8 membered aryl, cycloalkyl, heterocyclyl or heteroaryl ring, and

wherein each of R¹⁷ and R¹⁸ is independently selected from the group consisting of hydrido, halo, hydroxyl, alkoxy, amino, thio, sulfinyl, sulfonyl and

wherein R¹⁷ and R¹⁸ taken together can form a group consisting of ketal, thioketal,

$$= \begin{cases} = 0 , = \\ = s , = \\ = NOR^{22} \text{ and } = \end{cases} = NNR^{22}R^{23}$$

wherein each of R^{22} and R^{23} is independently selected from the group consisting of hydrido and alkyl.

3. A compound having the formula (I):

$$HO_2C$$
 HO_2C
 HO_2

and salts thereof;

wherein R is:

wherein X and X" are independently selected from C=O, C=S, C=NH, C=NR $^{\rm X}$, S=O or SO₂,

wherein n is 0 or 1;

wherein R^X is selected from alkyl, alkenyl, alkynyl, aryl, heteroaryl, cycloalkyl, heterocyclyl, hydroxyl, alkoxy, carboxy or carboalkoxy;

wherein B is X"RY, H, alkyl, alkenyl, alkynyl, aryl, heteroaryl, cycloalkyl or heterocyclyl,

wherein RY is selected from hydrido, alkyl, alkenyl, alkynyl, aryl, heteroaryl, cycloalkyl, heterocyclyl or hydroxyl;

wherein A is H, NH₂, NHR^A, NR^AR^B, alkyl, alkenyl, alkynyl, alkoxy, aryloxy, aryl, heteroaryl, cycloalkyl or heterocyclyl;

wherein R^A and R^B are independently selected from alkyl, alkenyl, alkynyl, aryl, heteroaryl, cycloalkyl, heterocyclyl or carboalkoxy;

wherein when n is 0, then A is additionally selected from:

wherein each of R⁵⁰-R⁵³ is independently selected from C₁-C₁₅ alkyl; alternatively, wherein B and A together form a 5-7 membered heterocyclic or heteroaryl ring;

wherein R¹ is

wherein X' and X"' are independently selected from C=0, C=S, C=NH, C=NR $^{X'}$, S=O or SO₂,

wherein m is 0 or 1;

wherein R^{X'} is selected from alkyl, alkenyl, alkynyl, aryl, heteroaryl, cycloalkyl, heterocyclyl, hydroxyl, alkoxy, carboxy or carboalkoxy;

wherein B' is X"'RY', H, alkyl, alkenyl, alkynyl, aryl, heteroaryl, cycloalkyl or heterocyclyl;

wherein R Y is selected from hydrido, alkyl, alkenyl, alkynyl, aryl, heteroaryl, cycloalkyl, heterocyclyl or hydroxyl;

wherein A' is alkyl, alkenyl, alkynyl, alkoxy or aryloxy, provided that when B' is H and X' is C=O, then A' is other than

- (a) -(C₁-C₁₆ unsubstituted alkyl)-NH₂;
- (b) $-(C_1-C_{10} \text{ unsubstituted alkyl})-NHC(O)R^D$, wherein R^D is $-C_1-C_{18}$ alkyl, optionally substituted with up to one hydroxyl, carboxyl or C_1-C_3 alkoxy, or one to three halo substituents;
- (c) $-C_1-C_{18}$ alkyl, optionally substituted with up to one hydroxyl, carboxyl or C_1-C_3 alkoxy, or one to three halo substituents;
 - (d) -C₄-C₁₈ unsubstituted alkenyl;

wherein R⁵⁴ is selected from C₁-C₁₇- unsubstituted alkyl or C₂-C₁₇- unsubstituted alkenyl; wherein R⁵⁵ is selected from hydroxyethyl, hydroxymethyl, mercaptomethyl, mercaptoethyl, methylthioethyl, 2-thienyl, 3-indolemthyl, phenyl optionally substituted with a group selected from halo, nitro, C₁-C₃-unsubstituted alkyl, hydroxy, C₁-C₃-unsubstituted alkoxy, C₁-C₃-unsubstituted alkylthio, carbamyl or C₁-C₃ unsubstituted alkylcarbamyl; or benzyl optionally substituted with a group selected from halo, nitro, C₁-C₃-unsubstituted alkyl, hydroxy, C₁-C₃-unsubstituted alkylcarbamyl; wherein t is 0 or 1 and wherein u is an integer from 1-3; and

when B is H and X is C=O, then X, together with A, does not form a carbamate amino protecting group; and

wherein when B' is H and m is 0, then A' is other than C₄-C₁₄ unsubstituted alkyl;

wherein R² is

wherein K and K' together form a C_3 - C_7 cycloalkyl or heterocyclyl ring or a C_5 - C_{10} aryl or heteroaryl ring;

wherein J is selected from the group consisting of hydrido, amino, NHR^J, NR^JR^K, alkyl, alkenyl, alkynyl, alkoxy, aryloxy, aryl, heteroaryl, cycloalkyl, heterocyclyl, alkylamino, hydroxyl, thio, alkylthio, alkenylthio, sulfinyl, sulfonyl, azido, cyano, halo,

$$- \begin{cases} S & \text{and} \\ - S & \text{oR}^{24} R^{25} \end{cases}$$

wherein each of R²⁴, R²⁵, and R²⁶ is independently selected from the group consisting of alkyl, cycloalkyl, heterocyclyl, aryl and heteroaryl; or R²⁴ and R²⁵ together form a 5-8 membered heterocyclyl ring;

wherein R^J and R^K are independently selected from alkyl, alkenyl, alkynyl, aryl, heteroaryl, cycloalkyl or heterocyclyl, or

alternatively, wherein J, together with R¹⁷, forms a 5-8 membered heterocyclyl or cycloalkyl ring, or

alternatively, wherein J, together with both R¹⁷ and R¹⁸, forms a 5-8 membered aryl, cycloalkyl, heterocyclyl or heteroaryl ring; and

wherein each of R¹⁷ and R¹⁸ is independently selected from the group consisting of hydrido, halo, hydroxyl, alkoxy, amino, thio, sulfinyl, sulfonyl and

wherein R^{17} and R^{18} taken together can form a group consisting of ketal, thioketal,

$$= \begin{cases} = 0 , = \\ = S , = \\ = NOR^{22} \text{ and } = \\ = NNR^{22}R^{23} \end{cases}$$

wherein each of R^{22} and R^{23} is independently selected from the group consisting of hydrido and alkyl.

4. A compound having the formula (I):

and salts thereof;

wherein R is:

wherein X and X" are independently selected from C=O, C=S, C=NH, C=NR $^{\rm X}$, S=O or SO₂,

wherein n is 0 or 1;

wherein R^X is selected from alkyl, alkenyl, alkynyl, aryl, heteroaryl, cycloalkyl, heterocyclyl, hydroxyl, alkoxy, carboxy or carboalkoxy,

wherein B is X"RY, H, alkyl, alkenyl, alkynyl, aryl, heteroaryl, cycloalkyl or heterocyclyl;

wherein RY is selected from hydrido, alkyl, alkenyl, alkynyl, aryl, heteroaryl, cycloalkyl, heterocyclyl or hydroxyl;

wherein A is H, NH₂, NHR^A, NR^AR^B, alkyl, alkenyl, alkynyl, alkoxy, aryloxy, aryl, heteroaryl, cycloalkyl or heterocyclyl,

wherein R^A and R^B are independently selected from alkyl, alkenyl, alkynyl, aryl, heteroaryl, cycloalkyl, heterocyclyl or carboalkoxy;

wherein when n is 0, then A is additionally selected from:

wherein each of R⁵⁰-R⁵³ is independently selected from C₁-C₁₅ alkyl; alternatively, wherein B and A together form a 5-7 membered heterocyclic or heteroaryl ring;

wherein R¹ is

wherein X' and X"' are independently selected from C=O, C=S, C=NH, C=NR $^{X'}$, S=O or SO₂,

wherein m is 0 or 1;

wherein R^{X'} is selected from alkyl, alkenyl, alkynyl, aryl, heteroaryl, cycloalkyl, heterocyclyl, hydroxyl, alkoxy, carboxy or carboalkoxy;

wherein B' and A' together form a 5-7 membered heterocyclic or heteroaryl ring;

wherein R^{A'} and R^{B'} are independently selected from alkyl, alkenyl, alkynyl, aryl, heteroaryl, cycloalkyl, heterocyclyl or carboalkoxy;

wherein R² is

wherein K and K' together form a C_3 - C_7 cycloalkyl or heterocyclyl ring or a C_5 - C_{10} aryl or heteroaryl ring;

wherein J is selected from the group consisting of hydrido, amino, NHR^J, NR^JR^K, alkyl, alkenyl, alkynyl, alkoxy, aryloxy, aryl, heteroaryl, cycloalkyl, heterocyclyl, alkylamino, hydroxyl, thio, alkylthio, alkenylthio, sulfinyl, sulfonyl, azido, cyano, halo,

$$- \begin{cases} S \\ NR^{24}R^{25} \end{cases} \text{ and } - \begin{cases} S \\ OR^{26} \end{cases}$$

wherein each of R²⁴, R²⁵, and R²⁶ is independently selected from the group consisting of alkyl, cycloalkyl, heterocyclyl, aryl and heteroaryl, or R²⁴ and R²⁵ together form a 5-8 membered heterocyclyl ring:

wherein R^{I} and R^{K} are independently selected from alkyl, alkenyl, alkynyl, aryl, heteroaryl, cycloalkyl or heterocyclyl; or

alternatively, wherein J, together with R^{17} , forms a 5-8 membered heterocyclyl or cycloalkyl ring; or

alternatively, wherein J, together with both R^{17} and R^{18} , forms a 5-8 membered aryl, cycloalkyl, heterocyclyl or heteroaryl ring; and

wherein each of R¹⁷ and R¹⁸ is independently selected from the group consisting of hydrido, halo, hydroxyl, alkoxy, amino, thio, sulfinyl, sulfonyl and

wherein R¹⁷ and R¹⁸ taken together can form a group consisting of ketal, thioketal,

$$= \begin{cases} = 0 & \text{if } = \\ = s & \text{if } = \\ = NOR^{22} \text{ and } = \\ = NNR^{22}R^{23} \end{cases}$$

wherein each of R²² and R²³ is independently selected from the group consisting of hydrido and alkyl.

5. The compound according to any of claims 1-4, wherein R is selected from the group consisting of:

wherein each of R^3 , R^4 R^5 , and R^6 is independently selected from the group consisting of hydrido, alkyl, aryl, heterocyclyl and heteroaryl, and wherein R^{44} is selected from the group consisting of alkyl, aryl, heterocyclyl and heteroaryl.

6. The compound according to claim 5, wherein R is selected from

$$R^{4}$$
 and R^{5} R^{4} R^{5}

wherein R^{4} is selected from the group consisting of alkyl, aryl-substituted alkyl, substituted phenyl, heterocyclyl, optionally substituted (C_8 - C_{14})-straight chain alkyl and SR^7 ; wherein R^7 is an alkyl group.

7. The compound according to claim 6, wherein R is selected from the group consisting of

$$R^3$$
 (C_8 - C_{13})-straight-chain alkyl (C_8 - C_{13})-straight-chain alkyl R^3 (C_8 - C_{13})-straight-chain alkyl R^3 and R^5 R^4 , R^5

wherein X^3 is chloro or trifluoromethyl and wherein q is 0 or 1.

8. The compound according to any of claims 1-4, wherein R¹ is selected from the group consisting of:

wherein R⁸ is selected from a natural amino acid side chain or an amino acid side chain that is not naturally occurring;

wherein each of R^9 , R^{10} and R^{11} is selected from hydrido, alkyl, aryl, heterocyclyl and heteroaryl;

wherein R^{12} is selected from the group consisiting of heterocyclyl, heteroaryl, aryl, and alkyl and

wherein R^{13} is selected from (C₁-C₃-alkyl) and aryl.

9. The compound according to claim 8, wherein R^1 is selected from the group consisting of:

$$R^{12}$$
, R^{8}

wherein R^8 is selected from tryptophan side chain and lysine side chain;

wherein each of R^{10} and R^{11} is independently selected from hydrido and alkyl;

wherein R^{12} is selected from imidazolyl, N-methylimidazolyl, indolyl, quinolinyl, benzyloxybenzyl, and benzylpiperidenylbenzyl; and wherein X is selected from fluoro, and trifluoromethyl.

10. The compound according to any of claims 1-4, wherein J is selected from the group consisting of hydrido, amino, azido and

wherein R^{17} and R^{18} taken together form a group selected from ketal,

$$= \begin{cases} = 0 & \text{and} & = \end{cases} = NOR^{22}$$

or wherein R¹⁷ is hydroxyl when R¹⁸ is hydrido; or wherein J, together with R¹⁷, forms a heterocyclyl ring.

11. The compound according to claim 10, wherein R^2 is selected from the group consisiting of

wherein R¹⁷ and R¹⁸ taken together form a group selected from

$$= \begin{cases} = 0 & \text{and} & = \end{cases} = NOR^{22}$$
, wherein R^{22} is selected from the group

consisting of H and alkyl, and wherein R¹⁹ is selected from the group consisting of

12. The compound according to claim 11, wherein R² is

13. The compound according to any one of claims 1-4 wherein said compound is selected from

Cpd #	· R	R ¹	R ²
1	NHCO(CH ₂) ₈ CH ₃	H NCO218U NHCO218U	O NH ₂
2	NHCO(CH ₂) ₈ CH ₃	.{-N	
3	NHCO(CH ₂) ₈ CH ₃	NHSO₂Ph	Ž-
4	NHCO(CH ₂) ₈ CH ₃	HN N N	Ž-
5	NHCO(CH ₂) ₈ CH ₃	HN H	- <u>z</u>
6	NHCO(CH ₂) ₈ CH ₃	HN H	Ž
7	NHCO(CH ₂) ₈ CH ₃	HN H	Ž-(-)
8	NHCO(CH ₂) ₈ CH ₃	S HN H	D T
9	NHCO(CH ₂) ₈ CH ₃	D N-NH HN N CO₂H	D =
10	NHCO(CH ₂) ₈ CH ₃	O NH ₂	\$ -\{-\}
11	NHCO(CH ₂) ₈ CH ₃	O NH2	NH ₂
. 12	NHCO(CH ₂) ₈ CH ₃	HN Br	NH2
13	NHCO(CH ₂) ₈ CH ₃	O NH ₂ CH ₃	O NH ₂
14	NHCO(CH ₂) ₈ CH ₃	O NH2 HN CH3	O NH2
15	NHCO(CH ₂) ₈ CH ₃	O NH ₂	O NH2
16	NHCO(CH ₂) ₈ CH ₃	O NH2 OCH3	O NH ₂

17	NHCO(CH ₂) ₈ CH ₃	HN CO	Ĭ-
18	NHCO(CH ₂) ₈ CH ₃	HN NO2	0 NH ₂
19	NHCO(CH ₂) ₈ CH ₃	O NH2 HN CO2H CO3H O NHCH3	O NH2
20	NHCO(CH ₂) ₈ CH ₃	HN	P. N. P.
21	NHCO(CH ₂) ₈ CH ₃	O OCH₃	ž -
22	NHCO(CH ₂) ₈ CH ₃	HN OH	NH ₂
23	NHCO(CH ₂) ₈ CH ₃	HN NH2	Ž-
24	NHCO(CH ₂) ₈ CH ₃	HN HA	O NH ₂
25	NHCO(CH ₂) ₈ CH ₃	HN PF	P. C.
26	NHCO(CH ₂) ₈ CH ₃	HN F.	O NH2
27	NHCO(CH ₂) ₈ CH ₃	HN F	NET -
28	NHCO(CH ₂) ₈ CH ₃	HN.	- E E
29	NHCO(CH ₂) ₈ CH ₃	HM	- N
30	NHCO(CH ₂) ₈ CH ₃	HN NH2	
31	NHCO(CH ₂) ₈ CH ₃	HŅ NH2	NH.
32	NHCO(CH ₂) ₈ CH ₃	O SCH ₃	o NH2

33	NHCO(CH ₂) ₈ CH ₃	HN	O NH.
34	NHCO(CH ₂) ₈ CH ₃	O N(CH ₃) ₂	O Ž
35	NHCO(CH ₂) ₈ CH ₃	HN	اب () ج
36	NHCO(CH ₂) ₈ CH ₃	O NHCH3	
37	NHCO(CH₂)8CH₃	HN S	22
38	NHCO(CH ₂) ₈ CH ₃	HN F	NH2
39	NHCO(CH ₂) ₈ CH ₃	о ососн ₃	O NH2
40	NHCO(CH ₂) ₈ CH ₃	HN OCH	O NH ₂
41	NHCO(CH ₂) ₈ CH ₃	HN NHBOC	O NH2
42	NHCO(CH ₂) ₈ CH ₃	HN CO₂CH₃	O NH2
43	NHCO(CH ₂) ₈ CH ₃	HN CO2'Bu	O NH2
44	NHCO(CH ₂) ₈ CH ₃	HN NHBOC N	NH ₂
45	NHCO(CH ₂) ₈ CH ₃	HN NH2	ZÉ,
46	NHCO(CH ₂) ₈ CH ₃	HN NH2 CO2CH3	O NH ₂
47	NHCO(CH ₂) ₈ CH ₃	O HN CONH₂ T NH₂	O NH ₂
48	NHCO(CH ₂) ₈ CH ₃	HN CONH ₂	O NH ₂
49	NHCO(CH ₂) ₈ CH ₃	O HN NHTS	O NH ₂

50	NHCO(CH ₂) ₈ CH ₃	HN	O NH ₂
51	NHCO(CH ₂) ₈ CH ₃	HŅ NH2 NHTS	O NH ₂
-52	NHCO(CH ₂) ₈ CH ₃	HN CH ₃	0 = \-\-\-\-\-\-\-\-\-\-\-\-\-\-\-\-\-\-\
54	NHCO(CH ₂) ₈ CH ₃	HN NH ₂	Ž
55	NHCO(CH ₂) ₈ CH ₃	HN NHBOC	
56	NHCO(CH ₂) ₈ CH ₃	HN NH ₂ OH	0=\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\
57	NHCO(CH₂)8CH₃	HN Cbz	Ž
58	NHCO(CH₂)8CH₃	HN S	O NH ₂
60	NHCO(CH ₂) ₈ CH ₃	HN NH2 NH2	
61	NHCO(CH₂)8CH₃	HY.	DHŽ
62	NHCO(CH ₂) ₈ CH ₃	HN NH, NH	O NH ₂
63	NHCO(CH₂)8CH₃	HAT NH2	O NH ₂
64	NHCO(CH ₂) ₈ CH ₃	NH, NH, NH	NH2
65	NHCO(CH ₂) ₈ CH ₃	HN NH ₂ O NH ₂	O NH ₂
66	NHCO(CH ₂) ₈ CH ₃	HN NH2 NH	O NH2
67	NHCO(CH ₂) ₈ CH ₃	HN NH ₂	O NH ₂

68	NHCO(CH ₂) ₈ CH ₃	HN NH ₂ S	0=\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\
69	NHCO(CH ₂) ₈ CH ₃	HN NH ₂ N N	ğ
70	NHCO(CH ₂) ₈ CH ₃	HN HN HN	- ZH
72	NHCO(CH ₂) ₈ CH ₃	HN NH2	ğ{-
73	NHCO(CH ₂) ₈ CH ₃	HN NH ₂ NH	\$
74	NHCO(CH ₂) ₈ CH ₃	HN NH ₂ NBOC	
75	NHCO(CH ₂) ₈ CH ₃	HN NHBOC	\$
76	NHCO(CH ₂) ₈ CH ₃	ÇH2-	ğ-{_}
77	NHCO(CH ₂) ₈ CH ₃	NH(CH ₂) ₂ OH	\$\frac{\frac}}}}}}}}{\frac{\frac{\frac{\frac{\frac{\frac{\frac{\frac{\frac{\frac}{\frac{\frac{\frac{\frac{\frac{\frac{\frac{\frac}{\frac}}}}}}}}}}}}}}}}}}}}}}}}}}}}}}}}}}}}
78	NHCO(CH ₂) ₈ CH ₃	HN N NHPh	O NH ₂
79	NHCO(CH ₂) ₈ CH ₃	NH NH	ğ
80	NHCO(CH ₂) ₈ CH ₃	NH N OCH3	D = T
81	NHCO(CH ₂) ₈ CH ₃	NH NH	0=
82	NHCO(CH ₂) ₈ CH ₃	NH CH ₃	\$ \frac{1}{2}
83	NHCO(CH ₂) ₈ CH ₃	HN CI	o NET?
84	NHCO(CH ₂) ₈ CH ₃	HN CH,	
85	NHCO(CH ₂) ₈ CH ₃	HN N	O NET

		,
HN NH ₂ NTs	HN NH ₂	NH ₂
NHCO(CH ₂) ₈ CH ₃	HN NO2	NET.
NHCO(CH ₂) ₈ CH ₃	N CI 2	O NH2
NHCO(CH ₂) ₈ CH ₃	HN CI	O = F
NHCO(CH ₂) ₈ CH ₃	N O O OMe2	O NH2
NHCO(CH ₂) ₈ CH ₃	N NE 12 2	O NH2
NHCO(CH ₂) ₈ CH ₃	HN NE 12	O NH ₂
NHCO(CH ₂) ₈ CH ₃	HN O Bu	O NH2
NHCO(CH ₂) ₈ CH ₃	HN O ⁿ Pr	O NH2
NHCO(CH ₂) ₈ CH ₃	HRY F	O NH2
NHCO(CH ₂) ₈ CH ₃	HN Meo	D N
NHCO(CH ₂) ₈ CH ₃	HN OME	NH2
NHCO(CH ₂) ₈ CH ₃	N F	0 N-
NHCO(CH ₂) ₈ CH ₃	N (F)	Ž -()
NHCO(CH ₂) ₈ CH ₃	HN O	O NH2
HN CI	HN NH ₂	O
NHCO(CH ₂) ₁₁ CH ₃	HN NH ₂	NH ₂
	NHCO(CH ₂) ₈ CH ₃	HN NH2 NHCO(CH2)8CH3 HN NH2 NHCO NHCO(CH2)8CH3 HN O O O O O O O O O O O O O O O O O O O

103	NHCO(CH ₂) ₈ CH ₃		O NH ₂
104	NHCO(CH ₂) ₈ CH ₃	YN NON O)2) =
105	NHCO(CH ₂) ₈ CH ₃	HN NO ₂	
106	NHCO(CH ₂) ₈ CH ₃	HN OH	N. A. C.
107	NHCO(CH ₂) ₈ CH ₃	HN CF	O NH2
108	NHCO(CH ₂) ₈ CH ₃	HN CI	
109	NHCO(CH ₂) ₈ CH ₃	HN CI	DHY
110	NHCO(CH ₂) ₈ CH ₃	N COO	DE T
111	NHCO(CH₂)8CH₃	HN	Ž-(-)
112	NHCO(CH ₂) ₈ CH ₃	N CF ₃	Ĭ- 0= -
113	NHCO(CH ₂) ₈ CH ₃	NE I	
114	NHCO(CH ₂)8CH₃	N NEI	
115	NHCO(CH ₂) ₈ CH ₃	HN HN	D Z
116	NHCO(CH ₂) ₈ CH ₃	HN CO	NH ₂
117	NHCO(CH ₂) ₈ CH ₃	HN O I'Bu	0 NH2

118	NHCO(CH ₂) ₈ CH ₃	N COOCI	O NH ₂
119	NHCO(CH ₂) ₈ CH ₃	HN CI	O NH2
120	NHCO(CH ₂) ₈ CH ₃	N NO2	O NH ₂
121	NHCO(CH ₂) ₈ CH ₃	HN 0	O NH,
122	NHCO(CH ₂) ₈ CH ₃	HŅ CO₂H	0 NH2
123	NHCO(CH ₂) ₈ CH ₃	HN O'Hex	0 NH2
124	NHCO(CH ₂) ₈ CH ₃	N O Hex	0 NH2
125	NHCO(CH ₂) ₈ CH ₃	N O'Bu	≥
126	NHCO(CH ₂) ₈ CH ₃	N O ⁿ Pr 2	\$\frac{\tau}{2}\\ 0=\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\
127	NHCO(CH ₂) ₈ CH ₃	× _{NH} —	
128	NHCO(CH ₂) ₈ CH ₃	N (0 - 0 F)	₹ - () o = () -
129	NHCO(CH ₂) ₈ CH ₃	HN NO2	DE T
130	NHCO(CH ₂) ₈ CH ₃	HN N	0=
131	NHCO(CH ₂) ₈ CH ₃	N (N)	NH.
132	NHCO(CH ₂) ₈ CH ₃	N OMe	

133	NHCO(CH ₂) ₈ CH ₃	HN NH2	O NH2
134	NHCO(CH ₂) ₈ CH ₃	N OMe	O NH ₂
135	NHCO(CH ₂) ₈ CH ₃	HN F	NH ₂
136	NHCO(CH ₂) ₈ CH ₃	HN	O NH ₂
137	NHCO(CH ₂) ₈ CH ₃	<u>"</u> (),	O NH2
138	NHCO(CH ₂) ₈ CH ₃	HN N	O NH2
139	NHCO(CH ₂) ₈ CH ₃	HN -	O NH2
140	NHCO(CH ₂) ₈ CH ₃		O NEZ
141	NHCO(CH ₂) ₈ CH ₃	N O D	O
142	NHCO(CH ₂) ₈ CH ₃	HN	
143	NHCO(CH ₂) ₈ CH ₃	HN CO	O NE 2
144	NHCO(CH ₂) ₈ CH ₃	HN Ph	O NHÃ
145	NHCO(CH₂)8CH₃	N Phy 2	O NH2
146	NHCO(CH ₂) ₈ CH ₃	Z Z Z Z Z Z Z Z Z Z Z Z Z Z Z Z Z Z Z	O NH ₂
147	NHCO(CH ₂) ₈ CH ₃	HN	O NH2
148	NHCO(CH ₂) ₈ CH ₃	m Z	O NH ₂

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NH ₂
O NH ₂

165	NHCO(CH ₂) ₈ CH ₃	N Q Ph)	O NH2
166	NHCO(CH₂)8CH₃	HN F	O NH2
167	NHCO(CH ₂) ₈ CH ₃	N()2	O NH2
168	NHCO(CH ₂) ₈ CH ₃	IN TO THE PART OF	O NH ₂
169	NHCO(CH₂)₃CH₃	N ()2	O NH2
171	NHCO(CH₂)8CH₃	N Butyl 2	O NH ₂
172	NHCO(CH ₂) ₈ CH ₃	HN nButyl	O NH2
173	NHCO(CH ₂) ₈ CH ₃	HN	O NH ₂
174	NHCO(CH ₂) ₈ CH ₃	HN S	£
175	NHCO(CH ₂) ₈ CH ₃	Pentyl	
176	NHCO(CH ₂) ₈ CH ₃	N T N	DE N
177	NH ₂		O NH ₂
178	NHCO(CH ₂) ₈ CH ₃	NH ₂	O NH ₂
179	NHCO(CH ₂) ₈ CH ₃	NHBOC NHBOC	O NH2
180	NHCO(CH ₂) ₈ CH ₃	HN NHF moc	O NH2
181	NHCONH(CH ₂) ₁₀ CH ₃	HN NH ₂	O NH2

182	. NHCO(CH ₂) ₈ CH ₃	HN	O NH2
183	NHCO(CH ₂) ₈ CH ₃	N OH)	NH,
184	NHCO(CH ₂) ₈ CH ₃	HNOH	O NHŽ
185	NHCO(CH ₂) ₈ CH ₃	HN CH	NH ₂
186	NHCO(CH ₂) ₈ CH ₃	N O O	O NH2
187	NHCO(CH₂)8CH₃	HN	NH2
189	NHCO(CH ₂) ₈ CH ₃	}-N	NH2
190	NHCO(CH ₂) ₈ CH ₃	O SO ₃ H	- NH.
192	NHCO(CH ₂) ₈ CH ₃	HN NH ₂ OH	
193	NHCO(CH ₂) ₈ CH ₃	B S S S S S S S S S S S S S S S S S S S	NH ₂
194	NHCO(CH ₂) ₈ CH ₃	HN OCF3	ZÉ.
195	NHCO(CH ₂) ₈ CH ₃	OCF ₃	NHZ T
196	NHCO(CH ₂) ₈ CH ₃	C - C	NH2
197	NHCO(CH ₂) ₈ CH ₃	HN CI	O NH2
198	NHCO(CH ₂) ₈ CH ₃	HN	NH2
199	NHCO(CH ₂) ₈ CH ₃	HN CI	NH ₂

200	NHCO(CH ₂) ₈ CH ₃	HN NH ₂ NH	+
201	NHCO(CH ₂) ₈ CH ₃	HN Hexyl	D T
202	NHCO(CH ₂) ₈ CH ₃	HN NMe ₂	O NH2
203	NHCO(CH ₂) ₈ CH ₃	HN NH 2 NH 2	DE C
204	NHCO(CH ₂) ₈ CH ₃	HN NHBo c N H	NH2
205	NH(CH ₂) ₈ CH ₃	HZ HZ	O H
206	NHCO(CH₂)8CO₂Me	O NH ₂	O = \
207	NHCO(CH ₂) ₆ CO ₂ Me	0 X X X X X X X X X	O NH2
208	NHCO(CH ₂) ₈ CH ₃	HN Ph	Ž.
209	NHCO(CH ₂) ₈ CH ₃	HN O F	O NH2
210	NHCO(CH ₂) ₈ CH ₃	HN CI	DEF2
211	NHCO(CH ₂) ₈ CH ₃	HN OBn	D = 1
212	NHCO(CH ₂) ₈ CH ₃	112 × 21	O NH ₂
213	NHCO(CH₂) ₆ NHBoc	NHBo c NH	O NH2
214	NHCO(CH ₂) ₇ NHBoc	O NHBo c NH	O NH2

215	NHCO(CH ₂) ₁₀ NHBoc	HN NHBo c	ž
216	NHCO(CH ₂) ₁₁ NHBoc	HN NHBo c NH	ž-(-)
217	NHCO(CH ₂) ₁₀ NH ₂	HN NH ₂ NH	
218	NHCO(CH ₂) ₁₁ NH ₂	HN NH ₂ N	D = -
219	NHCO(CH ₂) ₆ CH(CH ₃) ₂	HN NH ₂	
220	NHCONH(CH ₂) ₁₁ CH ₃	HN NH ₂ NH	
221	NHCO(CH ₂) ₈ CH ₃	HN NH N	£
222	NHCO(CH ₂) ₈ CH ₃		Ĭ
223	NHCO(CH₂)8CH₃	HIN N	Î -
224	NHCO(CH ₂) ₈ CH ₃	HN NHBo c	
225	NHCO(CH₂) ₈ CH ₃	HN NH ₂	£ ()
226	NHCO(CH ₂) ₈ CH ₃	HN N N	₹- - -
227	NHCO(CH₂)8CḤ3	HN N-Ph	
228	NHCO(CH ₂) ₈ CH ₃	HN N N F	£
229	NHCO(CH ₂) ₈ CH ₃	HN	ž-
230	NHCO(CH ₂) ₈ CH ₃	HN CI	₹

231	NHCO(CH ₂) ₈ CH ₃	HN Ph	
232	NHCO(CH ₂) ₈ CH ₃	HN N Ph	NH,
233	NHCO(CH ₂) ₈ CH ₃		Ž
234	NHCO(CH ₂) ₈ CH ₃		P P
235	NHCO(CH ₂) ₈ CH ₃	HN	Î-
236	NHCO(CH ₂) ₈ CH ₃		Ž
237	NHCO(CH ₂) ₈ CH ₃		Ž,
238	NHCO(CH ₂) ₈ CH ₃	HK	ž (
239	NHCO(CH ₂) ₈ CH ₃	HN N-Bn	Ž-\-
240	NHCO(CH ₂) ₈ CH ₃	HIN CO CO	\$
241	NHCO(CH ₂) ₈ CH ₃	HN N Ph	£-{}
242	NHCO(CH ₂) ₈ CH ₃		\$-\\-\\-\\-\\-\\-\\-\\-\\-\\-\\-\\-\\-\\
243	NHCO(CH ₂) ₈ CH ₃		\${\}
244	NHCO(CH ₂) ₈ CH ₃	HW C-C1	Ž-{-
245	NHCO(CH ₂) ₈ CH ₃	HN NH2	\$-\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\
246	HN CI	HN NH ₂ NH	O NHz
247	HN CI	HN NH ₂ N	D Z

248	HN OPh	HN NH ₂ NH	O NH2
249	HN O Buty	Hiv NH ² N	O NH2
250	HN C	HIN NH ₂ NH	O NH2
251	HN CI	HN NH ₂	NHŽ T
252	NHCO(CH ₂) ₈ CH ₃	HN NO ₂	O NH ₂
253	NHCO(CH ₂) ₈ CH ₃	HN N-Bn	O NH2
254	NHCO(CH ₂) ₇ CH ₃	NBœ HN NHBoc	NH2
255	NHCO(CH ₂) ₉ CH ₃	NBoc HN NHBoc	NH2
256	NHCO(CH ₂) ₁₀ CH ₃	NB C HN NHB oc	O NH ₂
257	NHCO(CH ₂) ₁₁ CH ₃	NB& HN_NHBoc	NH2
258	NHCO(CH ₂) ₁₂ CH ₃	NBoc HN NHBoc	NH2
259	NHCO(CH₂)8CH₃	NBbc HN NBbc	O NH ₂
260	NHCO(CH₂)₀CH₃	NH HN NH ₂	NH2
261	NHCO(CH ₂) ₁₁ CH ₃	NH HN NH ₂	O NH2
262	NHCO(CH ₂) ₁₂ CH ₃	NH HN NH ₂	O NH2

263	HN CI	NB& HN NHBoo	O NH ₂
264	HN N-uHebth	HN-NH ₂ NH ₂ NH	NH2
265	NHCO(CH ₂) ₈ CH ₃	HN CO	DE T
266	NHCO(CH ₂) ₈ CH ₃	HN NO ₂	DE NE
267	NHCO(CH ₂) ₈ CH ₃	HN NOZ	O NH2
268	NHCO(CH ₂) ₈ CH ₃	HN N-N-CF3	NH2
269	N=N N-heptyl	NHB% H	O NH2
270	NHCO(CH ₂) ₈ CH ₃	HN NH	DEP.
271	NHCO(CH ₂) ₈ CH ₃	HW CO	ŽŽ,
272	NHCO(CH ₂) ₈ CH ₃	N C C C C C C C C C C C C C C C C C C C	ž.
273	NHCO(CH ₂) ₈ CH ₃	N N OMe 2	NH2
274	NHCO(CH ₂) ₈ CH ₃	HN OMe	DH ₂
275	NHCO(CH ₂) ₈ CH ₃	HK LY CI	DE T
276	NHCO(CH ₂) ₈ CH ₃	N CI	NH2
277	NHCO(CH ₂) ₈ CH ₃	- OH	DE TO

278	NHCO(CH ₂) ₈ CH ₃	HN N	O NH2
279	NHCO(CH ₂) ₈ CH ₃	N N F	O NH ₂
280	NHCO(CH ₂) ₈ CH ₃	HIN CI	0 = \-\-\-\-\-\-\-\-\-\-\-\-\-\-\-\-\-\-\
281	NHCO(CH ₂) ₈ CH ₃		N-Y
282	NHCO(CH₂)₅CH₃	HN NO ₂	Ž
283	NHCO(CH ₂) ₈ CH ₃	HK C	D H
284	NHCO(CH ₂) ₈ CH ₃	N CI N CI	O NET Y
285	NHCO(CH ₂) ₈ CH ₃	H Z Z Z	≥±2
286	NHCO(CH ₂) ₈ CH ₃		DE N
287	NHCO(CH ₂) ₈ CH ₃		\$ = \frac{1}{1}
288	NHCO(CH ₂) ₈ CH ₃	HN CI OH	O NH2
289	NHCO(CH ₂) ₈ CH ₃	N(O4 ₅) ₂	O NH ₂
290	NHCO(CH ₂) ₈ CH ₃	+N N	Ž-
291	NHCO(CH ₂) ₈ CH ₃	DF CI	NH2

	γ		
292	HN CI	HN NH2	Ž.
293	NHCO(CH ₂) ₁₀ CH ₃	HIV NH2	O NE
294	NHCO(CH ₂) ₇ CH ₃	NH HN NH ₂	O NET ?
295	NHCO(CH ₂) ₁₁ CH ₃	HN NHBoc	O NH2
296	NHCO(CH ₂) ₁₀ CH ₃	NH Boc NH Boc	NHÎ -
297	NHCO(CH ₂) ₉ CH ₃	O HN NHBoc NHBoc	P. T.
298	NHCONH(CH₂)7CH₃	HN NHBoc NHBoc	D NE Î
299	NHCONH(CH ₂) ₁₀ CH ₃	HN NHBoc NH Boc	NHŽ T
300	NHCONH(CH ₂) ₁₁ CH ₃	HN NH Boc	PE -
301	NHCO(CH ₂) ₁₁ CH ₃	HN NH ₂	→ NH ₂
302	NHCO(CH ₂) ₁₀ CH ₃	HN NH ₂	NH.
303	NHCO(CH ₂) ₉ CH ₃	O HN NH ₂	PÉ -
304	NHCONH(CH₂)7CH₃	NH ₂	P. P
305	NHCONH(CH ₂) ₁₀ CH ₃	HN NH ₂	O NH ₂
306	NHCONH(CH ₂) ₁₁ CH ₃	HN NH ₂	O NH2
307	NHCO(CH ₂) ₉ CH ₃	HN NHBoc N H	o NET

308	NHCO(CH ₂) ₁₀ CH ₃	NHB% ZH	NH2
309	NHCO(CH ₂) ₁₀ CH ₃	HN-NH ₂	O NET?
310	NHCO(CH ₂) ₉ CH ₃	HN NH ₂	O NET?
311	NHCONH(CH₂), CH₃	HN NH2 NH2 NH2 NH2 NH2 NH2 NH2 NH2 NH2 N	O NH2
312	NHCONH(CH₂)7CH₃	HN F	NE?
313	NHCONH(CH ₂) ₇ CH ₃	HN NH ₂	NH ₂
314	NHCONH(CH ₂) ₁₀ CH ₃	NBoc HN NHBoc	PÉ -
315	NHCONH(CH₂)⁊CH₃	HN OCH	O NH ₂
316	NHCONH(CH ₂) ₇ CH ₃	HN HN	O NH2
317	NHCONH(CH ₂) ₇ CH ₃	HN NO ₂	
318	NHCO(CH ₂) ₉ CH ₃	HN N H	O NEZ
319	NHCO(CH ₂) ₉ CH ₃	HN HN	
320	NHCO(CH ₂) ₁₁ CH ₃	HN OCH3	O NH ₂
321	NHCO(CH ₂) ₁₁ CH ₃	HN NO ₂	Σ- - - -
322	NHCO(CH ₂) ₁₁ CH ₃	HN N	O NH ₂

323	NHCO(CH ₂) ₈ CH ₃	HN CF3	O NH2
324	NHCO(CH ₂) ₈ CH ₃	HN CF ₃	
325	NHCO(CH ₂) ₈ CH ₃	HN F	0 - \-\-\-\-\-\-\-\-\-\-\-\-\-\-\-\-\-\-
326	NHCO(CH ₂) ₈ CH ₃	HN F	D =
327	NHCO(CH₂)8CH₃	HN F	Ž
328	NHCO(CH ₂) ₈ CH ₃	HN CF ₃	€-{-}
329	NHCO(CH ₂) ₈ CH ₃	HN CF ₃	O NH2
330	NHCO(CH ₂) ₈ CH ₃	HN CI CI F	O NH2
331	NHCO(CH ₂) ₈ CH ₃	HN CF ₃	\$ -\\ -\\ -\\ -\\ -\\ -\\ -\\ -\\ -\\ -\
332	NHCO(CH ₂) ₁₀ CH ₃	HN OCH3	- - - - -
333	NHCO(CH ₂) ₁₀ CH ₃	HN NO2	ğ
334	NHCO(CH ₂) ₁₀ CH ₃	HN N	NH.
335	NHCONH(CH ₂) ₁₁ CH ₃	HN HN	- \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \
336	NHCONH(CH ₂) ₁₁ CH ₃	HŅ NO ₂	\$\frac{1}{2}
337	NHCONH(CH ₂) ₁₁ CH ₃	HN N H	0 NH2
338	NHCO(CH ₂) ₁₂ CH ₃	HN NHBoc N H	≥ ₹2

339	NHCO(CH ₂) ₁₂ CH ₃	HN NH2 N	O NH2
340	NHCO(CH ₂) ₁₂ CH ₃	NHBo c	O NH ₂
341	NHCO(CH ₂) ₁₂ CH ₃	HN NH ₂	O NH2
342	NHCO(CH ₂) ₉ CH ₃	HN P	O NH ₂
343	NHCO(CH ₂) ₁₀ CH ₃	HN HN NH2	DE T
344	NHCO(CH ₂) ₁₂ CH ₃	HN F HN	₹
345	NHCO(CH ₂) ₁₂ CH ₃	HN HN	£
346	NHCO(CH₂)₁₂CH₃	HN OCH,	DE T
347	NHCO(CH ₂) ₇ CH ₃	HN NH	o ₹2
348	NHCO(CH ₂) ₇ CH ₃	HN HN	O NH2
349	NHCO(CH ₂) ₇ CH ₃	HN NO ₂	ž
350	HN CI	HN F NH2	NE Î
351	NHCO(CH ₂) ₁₁ CH ₃	HN PH2	NH2
352	NHCONH(CH ₂) ₁₀ CH ₃	HN + 1	o NET
355	NHCONH(CH ₂) ₁₀ CH ₃	HN HN	O NH ₂
356	NHCONH(CH ₂) ₁₀ CH ₃	HN N H	O NH2

	T		
358	NHCO(CH ₂) ₈ CH ₃	HN O H	NH ₂
359	NHCO(CH ₂) ₈ CH ₃	HN 0 H - 0 H - 0 N N N N N N N N N N N N N N N N N N	O NH2
360	NHCO(CH ₂) ₈ CH ₃	HN S-N NCH	O NH.
361	NHCO(CH ₂) ₈ CH ₃	HN OH,	O NE
362	NHCO(CH ₂) ₈ CH ₃	HN S-N NPh	O NH2
363	NHCO(CH ₂) ₈ CH ₃	HN - S-N N-N	O NH ₂
364	NHCO(CH ₂) ₈ CH ₃	0 =5-N -0 -0 -0 -0 -0 -0 -0 -0 -0 -0 -0 -0 -0	Ĕ- -
365	NHCO(CH₂)8CH₃	HN OS NBO	O NH ₂
366	NHCO(CH ₂) ₈ CH ₃		O NH2
367	NHCO(CH ₂) ₈ CH ₃	HN S-N N-N N-N N-N N-N N-N N-N N-N N-N N-	O NH2
368	NHCO(CH ₂) ₈ CH ₃	HM - S-N N - CI	NH ₂
369	NHCO(CH ₂) ₈ CH ₃	HIX	NH2
370	NHCO(CH ₂) ₈ CH ₃	HN - 5-N N- 5-F	ZH2
371	NHCO(CH ₂) ₈ CH ₃	HN	O NH ₂
372	NHCO(CH ₂) ₈ CH ₃	HN - 5-N N- C5	O NH2
373	NHCO(CH ₂) ₈ CH ₃	HN	O NH ₂
374	NHCO(CH ₂) ₈ CH ₃	HN N - N - N - N - N - N - N -	O NH ₂

375	NHCO(CH ₂) ₈ CH ₃	HN S-N F	O NH ₂
376	NHCO(CH ₂) ₈ CH ₃	HN O H	O NH ₂
377	NHCO(CH ₂) ₈ CH ₃	HN	NH ₂
378	NHCO(CH ₂) ₈ CH ₃	HIX 0 1 5 5 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7	ğ(-)
379	NHCO(CH ₂) ₈ CH ₃	HĀ O	Ž
380	NHCO(CH ₂) ₈ CH ₃	0 = 5 - C	Ž-(-)
381	NHCO(CH ₂) ₈ CH ₃	0:5-7 F	Ž
382	NHCO(CH ₂) ₈ CH ₃	HV	
383	NHCO(CH ₂) ₈ CH ₃	HN	
384	NHCO(CH ₂) ₈ CH ₃	HZ	
385	NHCO(CH ₂) ₈ CH ₃	9 H 5 N S N S N S N S N S N S N S N S N S N	O NH2
386	NHCO(CH ₂) ₈ CH ₃	HN 0 H CF3	O NH2
387	NHCO(CH ₂) ₈ CH ₃	нм — — — — — — — — — — — — — — — — — — —	O NH2
388	NHCO(CH ₂) ₈ CH ₃	HN	O NH ₂
389	NHCO(CH ₂) ₈ CH ₃	HN	O NH ₂
390	NHCO(CH ₂) ₈ CH ₃	HN CI	O NH2
391	NHCO(CH ₂) ₈ CH ₃	HN CI	O NH ₂

392	NHCO(CH ₂) ₈ CH ₃	المناسبة الم	O NH ₂
393	· NHCO(CH ₂) ₈ CH ₃		O NH ₂
394	NHCO(CH ₂) ₈ CH ₃	HK 0.0 0.3 0.3 0.4 0.4 0.4 0.4 0.4 0.4 0.4 0.4 0.4 0.4	D NH2
395	NHCO(CH₂)8CH₃	HK	Ĭ- 0= {-
398	NHCO(CH₂)8CH₃	HN TN F	Ž
399	NHCO(CH ₂) ₈ CH ₃	HN	NH2
400	NHCO(CH ₂) ₈ CH ₃	HŅ N	D N N N N N N N N N N N N N N N N N N N
401	NHCO(CH₂)8CH₃	HN N N N N N N N N N N N N N N N N N N	Property of the state of the st
402	NHCO(CH ₂) ₈ CH ₃	HN N S N N N N N N N N N N N N N N N N N	NH2
403	NHCO(CH ₂) ₈ CH ₃	HZ 2 50 50 50 50 50 50 50 50 50 50 50 50 50	
404	NHCO(CH ₂) ₈ CH ₃	HN CF3	O NH2
405	NHCO(CH ₂) ₈ CH ₃	Z , , , , , , , , , , , , , , , , , , ,	NH2

406	NHCO(CH ₂) ₈ CH ₃	T Z Z	€
407	NHCO(CH ₂) ₈ CH ₃	HN CO	
408	NHCO(CH ₂) ₈ CH ₃	HN CI	DE STATE OF THE ST
409	NHCO(CH ₂) ₈ CH ₃	HN	DHŽ
410	NHCO(CH ₂) ₈ CH ₃	, NH	O NH ₂

14. The compound according to claim 13 selected from the group consisting of

Cpd #	R	R ¹	R ²
45	NHCO(CH₂)8CH₃	HN NH ₂	NH,
54	NHCO(CH₂)8CH₃	HN NH ₂	O NET
76	NHCO(CH ₂) ₈ CH ₃	CH ₂ —	\$
81	NHCO(CH₂)8CH₃	NH NH	O T
85	NHCO(CH₂)8CH₃	HN N	0 H
102	NHCO(CH₂)11CH₃	HN NH2 NH	
209	NHCO(CH ₂) ₈ CH ₃	HN O F	0 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1
212	NHCO(CH ₂) ₈ CH ₃	HN Z	0 1
253	NHCO(CH ₂) ₈ CH ₃	HIV V-Bn	\$
260	NHCO(CH ₂) ₉ CH ₃	NH HN NH ₂	NH2
262	NHCO(CH ₂) ₁₂ CH ₃	NH HN NH ₂	O NH2
282	NHCO(CH ₂) ₈ CH ₃	HN NO ₂	NH2
285	NHCO(CH ₂) ₈ CH ₃	HX CI	O NH2

319	NHCO(CH ₂) ₉ CH ₃	HN HN	€
322	NHCO(CH ₂) ₁₁ CH ₃	HN HN	O NH2
333	NHCO(CH ₂) ₁₀ CH ₃	HŅ NO2	0 NH2
334	NHCO(CH ₂) ₁₀ CH ₃	HN HN	\$ 1 × 1
335	NHCONH(CH₂)11CH₃	HN HN	\$ 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1
336	NHCONH(CH ₂) ₁₁ CH ₃	HN NO2	NIT'S
344	NHCO(CH ₂) ₁₂ CH ₃	O NH ₂	O NH
355	NHCONH(CH ₂) ₁₀ CH ₃	HN HN	0 NH ₂

15. A compound of formula (I) according to claim 1, wherein R is NHCO-[(C_6 - C_{14})-alkyl]-CH₃, and R¹ and R² are selected from:

R	R ²
-}-N NCO₂IBu	O NH ₂
NHCO2tBu	
-}-N \ NH	O NH ₂
NH ₂	+ 0
NHSO₂Ph	- NET
HN TH	O NH2
#	N. S.
HN HN	O NH?
HN H	O NH,
HN H	O NH ₂
N-NH CO ₂ H	O NH ₂
O NH ₂	O NH ₂
O NH ₂	O NH
O NH ₂ HN HN O NH ₂	NE?
HN CH ₃	O NET
O NH ₂ HN CH ₃ O NH ₂	O NH2
HN OCH3	O NH.
HN OCH3	O NH2

HN CI	NH;
N ₃	NH ₂
ну Соэн	NH2
O NHCH ₃	O NH.
HN OCH,	O NH.
O NH ₂	O NH2
HŅ NH.	O NH ₂
O NH ₂	O NH ₂
HN F	DE LE
O NH ₂	O NH2
O NH ₂	O NH2
HN	O NH ₂
HN N	O NH ₂
HŅ NH2	O NH2
O HN NH ₂	O NH ₂
o sch	O NH ₂

HN	O NH,
O N(CH ₃) ₂	O NH ₂
HN	O NH2
HN NHCH,	+
HN NH	
HN F	O NH2
0 0COCH₃ HN CO2H	O NH.
HN OCH3	O NH2
HN NHBOC	O NH ₂
O HN CO ₂ CH ₃	NH.
HN CO2'Bu NHBOC	O NH2
HN NHBOC N	O NH ₂
HN NH ₂ NH	O NH2
O CO ₂ CH ₃	O NH ₂
HN CONH ₂	O NH ₂
HN CONH,	O NH ₂
O HN NHBOC	NH ₂

HN N	NH ₂
HN NH ₂ NHTs	O NH ₂
HN CH3	NH ₂
HN NH ₂	NH2
ни ннвос	O NH2
HN NH ₂ OH	O NH2
HN Cbz	O NH ₂
HN NH	NH2
HN NH ₂ NH ₂	NH2
HN T	NH ₂
HN NH ₂ NH	NH ₂
HN NH ₂	O NH ₂
HIV NH, NH	O NH2
HN NH ₂ O NH ₂	O NH ₂
HN NH2 H	O NH ₂
HN NH ₂	O NH2

HN	O NIT'S
NH ₂ S	+ 0
HŅ	O NH ₂
NH₂ N≈/	1
HN	NH,
Н	+
HŅ NH2	O NH ₂
0 	O NH2
HN	
NH ₂ N = /NH ₂	O NH ₂
HN NH ₂ NBOC	
0	O NH ₂
HN NHBOC	
	O NH ₂
NH(CH ₂)₂OH	-
HN N NHPh	O NH ₂
NH ₂	+0
NH (O NH ₂
, N	+ 0
NH OCH,	O NH ₂
	+
NH F	O NH ₂
T 'N	+ -
NH N	O NH ₂
СH ₃	
HN O	NH ₂
c/~	+ -
HN N	O NH ₂
	+ -
HM (N)	O NH ₂
N-J	O NH ₂
HN NO2	O NH ₂
<u> </u>	

"(C C C)	O NH
HN CI	NH ₂
N O O OME	O NH2
N NE 12	O NH,
HN NE 12	NH ₂
HIN O'BU	O NH2
HN O Pr	O NH2
HN OFF	O NH _z
HN	O NH.
OMe HN C	O NH ₂
N ()	O NH ₂
N F	NH2
HN	O NH2
YN NON	O NH2
×	O NH ₂
HN NO	O NH ₂

HN OH	NE Z
HN O CF	O NH;
HN CI) 0 1
HN CI	E
N COO	-} -} - <u>\$</u>
HK CO	DH.
N CF3	D = 1
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N NEI	ž -
HN NC	ž.
HN	ğ
HN O O	O NE
N Co Co Co 2	0 = 1 = 1 = 1 = 1 = 1 = 1 = 1 = 1 = 1 =
HN CI	O NH ₂
N (0 NO ₂)	\$ -\-\-
HN~~~	O NH

HN CO₂H	O NH ₂
HN O'Hex	O NH2
N OnHex	O NH?
N O'Bu 2	O NH,
N O'Pr	
NH NH	O NH ₂
N CO CO F	
HN NO2	£
HN N	O NH2
N (N)	D Z Z
OMe 1	O NH2
HN H2	O NH ₂
N OMe	O NH ₂
HN F	O NH ₂
HN	O NH2

M O	
	NH,
HA T	NH2
N CO	o Fr
N O D	Ž -
HN O	
HN O O	£-(_)
HIV CO	\$
N PP) 2	0 ZH2
TZ Z Z	0 NH2
HILL COLOR	NH2
	O NH ₂
HIN O N	O NH ₂
Ph HN Z N H	O NH2
HIN OME	O NH2
N OMe O	O NH ₂

HN O'Dodecyl	NH2
HN O ⁿ De cyl	O NH2
HN O ⁿ Octyl	O NH2
HN CO ₂ H	O NH ₂
HN NMb2	O NH2
HN S	O NH ₂
HN N-Ph	O NH2
N Ph	O NH2
HN CO2H	O NH ₂
N - B) - F	O NH ₂
N NO2/2	O NH ₂
HIN OOD TO	O NH ₂
N Q Ph	O NH,
HN F	O NH ₂
N ()	NH?
HN	O NH ₂

N ()2	O NH2
N ngulyi	O NH ₂
HN Butyl	DEF2
HN	NH,
HN S	NH,
HN	\$ -\{-\}
N	0 N N
## \\	\$
HN NH ₂	\$ - \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \
NH ₂	D E
NHF moc	£
HN III	O EE
N OH)	
HN	£
HN OH) 0=
N O O	O NH2

H. H.	NH2
)-n	0 NH ₂
O SO3H	O NH ₂
HN NH ₂ OH	+ 2
Boc HN	O NH ₂
HN OCF3	±
N OCF ₃	DE ZE
ō → Ç	ž O
HN CI	Ē
HN N(CH ₃) ₂	₹ 0= {-
CT PT CT	P. P.
HN NH ₂ NH	+
HN nHexyl	O NH ₂
HN NMey 2	O NH ₂
HN NH ₂ NH ₁	O NH ₂
HN NHBoc NH	O NH ₂

	O NH2
HN SO	N. T.
HN O F	0
HN CI	O NH ₂
HN OBn	NH2
HZ Z ZH	DH.
H Z Z Z Z Z Z Z Z Z Z Z Z Z Z Z Z Z Z Z	O NH2
	N. P.
HN N	PE T
HN NHBo c	D H
HN NH ₂	O NH ₂
HIT ON NOT OF	O NH ₂
HN N-Ph	O NH2
HN N- F	O NH ₂
HK STONE	O NH2
HN CI	O NH ₂
HN N Ph	O NH2
HN Ph	NH2

HN - N N N	\$
HN - N - N - N - N - N - N - N - N - N -	O NEZ
HIN N-C-CI	O NH2
	O NH ₂
##\	O NH2
HK	Î -
HN N-Bn	ğ
HIV C1	O NH ₂
HIN Ph	NH ₂
	O H
	O NH ₂
HIV > - C1	O NH ₂
HN NH ₂	O NH,
O NH ₂ NH ₂	O Hĩ
HN NO, N-Bn	O NH2
HN N-Bn	O NH2
NBbc HN NBbc	O NH2
CI C	O NH2

	O NH ₂
HN NO ₂	+0
HN S	D Z Z
HIN NON-CE	O N
NH HN NH	O NH.
HIN CI	O NH ₂
N C C C C C C C C C C C C C C C C C C C	O NH,
N OMe	D E Î
HN N OMe	Î-
HN CI	Ē -
N CI)	O NH2
HIN OH	O NH2
HIN N	O NH ₂
N N N	O NH ₂
HM CI	O NH ₂
N Cl	O NH ₂
HN NO ₂	NH.

ÇI	O NH ₂
HIV THE	+
N CI	\$\tilde{\text{F}}
HZ CI	NH.
-\frac{1}{2}	0 2 1 2
HZ A	
HE COH	₹
M(CH ₂) ₂	E
HN N	NH ₂
OH CI	≥ ₹2
NH Boc	DE C
HN NH OCH3	O NH2
HN NO ₂	- T
HN CF3	Ê -
HN CF ₃	o z z

	A ARL
HN F	\$ - \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \
HN F	≥ ± 1
HN TO	O NH ₂
HN CI CF3	0 NH ₂
HN CF3	NH ₂
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16. The compound according to claim 15, wherein R is select d from NHCO-[(CH₂)₆₋₁₄]-CH₃.

- 17. A pharmaceutical composition comprising the compound according to any one of claims 1-4 and a pharmaceutically acceptable carrier.
- 18. A method of treating a bacterial infection in a subject, comprising the step of administering a therapeutically-effective amount of the pharmaceutical composition according to claim 17 to a subject in need thereof.
- 19. The method according to claim 18, wherein said subject is selected from the group consisting of a human, an animal, a cell culture or a plant.
- 20. The method according to claim 18, wherein said bacterial infection is caused by a gram-positive bacteria.
- 21. The method according to claim 20, wherein said bacteria is an antibiotic-resistant bacteria.
- 22. The method according to claim 21, wherein said antibiotic-resistant bacteria are resistant to an antibiotic selected from the group consisting of vancomycin, methicillin, glycopeptide antibiotics, penicillin and daptomycin.
- 23 The method according to claim 18, further comprising the step of co-administering more than one compound of Formula (I) to a subject in need thereof.
- 24. The method according to claim 18, further comprising the step of co-administering an antimicrobial agent other than a compound of Formula (I) to a subject in need thereof.

25. The method according to claim 22, wherein said antimicrobial agent is selected from the group consisting of penicillins and related drugs, carbapenems, cephalosporins and related drugs, aminoglycosides, bacitracin, gramicidin, mupirocin, chloramphenicol, thiamphenicol, fusidate sodium, lincomycin, clindamycin, macrolides, novobiocin, polymyxins, rifamycins, spectinomycin, tetracyclines, vancomycin, teicoplanin, streptogramins, anti-folate agents including sulfonamides, trimethoprim and its combinations and pyrimethamine, synthetic antibacterials including nitrofurans, methenamine mandelate and methenamine hippurate, nitroimidazoles, quinolones, fluoroquinolones, isoniazid, ethambutol, pyrazinamide, para-aminosalicylic acid (PAS), cycloserine, capreomycin, ethionamide, prothionamide, thiacetazone, viomycin, eveminomycin, glycopeptide, glycylcylcline, ketolides, oxazolidinone; imipenen, amikacin, netilmicin, fosfomycin, gentamicin, ceftriaxone, Ziracin, LY 333328, CL 331002, HMR 3647, Linezolid, Synercid, Aztreonam, and Metronidazole, Epiroprim, OCA-983, GV-143253, Sanfetrinem sodium, CS-834, Biapenem, A-99058.1, A-165600, A-179796, KA 159, Dynemicin A, DX8739, DU 6681; Cefluprenam, ER 35786, Cefoselis, Sanfetrinem celexetil, HGP-31, Cefpirome, HMR-3647, RU-59863, Mersacidin, KP 736, Rifalazil; Kosan, AM 1732, MEN 10700, Lenapenem, BO 2502A, NE-1530, PR 39, K130, OPC 20000, OPC 2045, Veneprim, PD 138312, PD 140248, CP 111905, Sulopenem, ritipenam acoxyl, RO-65-5788, Cyclothialidine, Sch-40832, SEP-132613, micacocidin A, SB-275833, SR-15402, SUN A0026, TOC 39, carumonam, Cefozopran, Cefetamet pivoxil, and T 3811.

- 26. The method according to claim 22, wherein said antimicrobial agent is selected from the group consisting of imipenen, amikacin, netilmicin, fosfomycin, gentamicin, ceftriaxone, teicoplanin, Ziracin, LY333328, CL331022, HMR3647, Linezolid, Synercid, Aztreonam and Metronidazole.
- 27. The method according to claim 19, wherein said subject is selected a human or an animal.

28. The method according to claim 27, wherein said subject is a

29. A compound having the formula (III):

human.

wherein R¹⁵ is selected from hydrido and an carbamate amino protecting group, preferably a *tert*-butoxycarbonyl group, wherein R¹⁶ is selected from the group consisting of

wherein R⁵⁷ is a halo or halo substituted alkyl group, preferably a fluoro or trifluoromethyl group; wherein, R²⁰ is an amino acid side chain, preferably a lysine or tryptophan side chain.

30. The compound according to claim 29 selected from:

Compound #	R ¹⁶
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25	O NH ₂
45	H ₂ N N
54	NH ₂
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80	OCH ₃
81	-\-\-\-\-\-\-\-\-\-\-\-\-\-\-\-\-\-\-\
82	- 12 N
84	- N-N-N-N-N-N-N-N-N-N-N-N-N-N-N-N-N-N-N

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174 78 78 150 130 138 168 274 274 317		7. H
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150 NH ₂ Ph 130 138 168 274 274 317		, \(\frac{1}{2} \) \(\frac{1} \) \(\frac{1} \) \(\frac{1}{2} \) \(\frac{1}{2} \
130 138 168 274 274 317	78	
138 168 274 274 317 317	150	
138 168 274 274 317 317		N H
274 274 317 N OCH ₃	130	- \(\)
274 274 317 N OCH ₃		\\
317. CCH ₃	168	-74. N
317. " " " N N	-	
	317.	NO ₂

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164	, '\(\)
210	NO ₂
107	CF ₃

111	- 1/2 O O
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253	
227	N. Ph
372	CF ₃
386	S CF3

31. A method of using the compound according to either of claims 29 or 30 to make a compound according to any one of claims 1-4.

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A. CLASSIFICATION OF SUBJECT MATTER IPC 7 C07K7/08 C12R C12R1/465 According to International Patent Classification (IPC) or to both national classification and IPC **B. FIELDS SEARCHED** Minimum documentation searched (classification system followed by classification symbols) IPC 7 C07K C12R Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched Electronic data base consulted during the international search (name of data base and, where practical, search terms used) EPO-Internal, WPI Data, CHEM ABS Data, BIOSIS, EMBASE C. DOCUMENTS CONSIDERED TO BE RELEVANT Relevant to claim No. Citation of document, with indication, where appropriate, of the relevant passages 1,3-5,29X EP 0 095 295 A (LILLY CO ELI) 30 November 1983 (1983-11-30) 1 - 31claim 1; tables 1-18 EP 0 178 152 A (LILLY CO ELI) 1,3,4,29 X 16 April 1986 (1986-04-16) claim 1 1-31 US 4 537 717 A (DEBONO MANUEL ET AL) 1,3,4,29 27 August 1985 (1985-08-27) 1-31 Υ claim 1 Further documents are listed in the continuation of box C. Patent family members are listed in annex. Special categories of cited documents: "T" later document published after the international filing date or priority date and not in conflict with the application but "A" document defining the general state of the art which is not cited to understand the principle or theory underlying the considered to be of particular relevance invention "E" earlier document but published on or after the international "X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone *L* document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified) "Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such docu *O* document referring to an oral disclosure, use, exhibition or ments, such combination being obvious to a person skilled in the art. other means document published prior to the international filing date but "&" document member of the same patent family later than the priority date claimed Date of the actual completion of the international search Date of mailing of the international search report 22 May 2001 30/05/2001 Authorized officer Name and mailing address of the ISA European Patent Office, P.B. 5818 Patentlaan 2 NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo nl, Fax: (+31-70) 340-3016 Deffner, C-A

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